APS Search for 09/123,620 FILE 'USPAT' ENTERED AT 12:21:12 ON 22 MAY 1999

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E#	FILE	FREQUENCY TERM
Εl	USPAT	I ELFMARK, JIRI/IN
E2	USPAT	3 ELFNER, BO A/IN
E3	USPAT	0> ELFORD H/IN
E4	USPAT	i ELFORD, ANDREW M/IN
E5	USPAT	3 ELFORD, DAVID/IN
E6	USPAT	8 ELFORD, HOWARD L/IN
E7	USPAT	3 ELFORD, PETER ELLICE/IN
E8	USPAT	1 ELFORD, WILLIAM J/IN
E9	USPAT	1 ELFRING, GARY C/IN
E10	USPAT	4 ELFSTRAND, JAMES K/IN
EH	USPAT	1 ELFSTRAND, STIG OLOF/IN
E12	USPAT	2 ELFSTROM, BO/IN

8 *ELFORD, HOWARD L*/IN LI

=> d 1-8 bib ab

US PAT NO: 5,366,996 [IMAGE AVAILABLE] DATE ISSUED: Nov. 22, 1994 Method of treating hemoglobinopathies TITLE: INVENTOR: **Howard L. Elford**, 3313 Gloucester Rd., Richmond, VA 23227 Bartholomeus van't Riet, 3419 Noble Ave., Richmond, VA 23222 APPL-NO: 07/986,861 DATE FILED: Dec. 7, 1992 ART-UNIT: 125 PRIM-EXMR: Marianne M. Cintins ASST-EXMR: M. Moezie LEGAL-REP: James L. Rowe

US PAT NO: 5,366,996 [IMAGE AVAILABLE] L1: 1 of 8

L1: 1 of 8

L1: 2 of 8

A therapeutic process for treating anemias in primates, including man, particularly those anemias of genetic origin including sickle-cell anemia, which comprises administering to an anemic primate an amount of a polyhydroxy benzoic, mandelic or phenylacetic acid derivative as specified at a dose level sufficient to increase fetal hemoglobin.

US PAT NO: 5,350,770 [IMAGE AVAILABLE] DATE ISSUED: Sep. 27, 1994

Therapeutic process for the treatment of septic shock

INVENTOR: **Howard L. Elford**, 3313 Gloucester Rd., Richmond, VA 23227

Bartholomeus van't Riet, 3419 Noble Ave., Richmond, VA 23222

APPL-NO: 07/919,907 DATE FILED: Jul. 28, 1992 PRIM-EXMR: Mo

Marianne M. Cintins William R. A. Jarvis LEGAL-REP: James L. Rowe

US PAT NO: 5,350,770 [IMAGE AVAILABLE] L1: 2 of 8

ABSTRACT:

A therapeutic process for treating septic shock comprising the administration of a polyhydroxy-substituted benzamide or phenylacetamide derivative to a human suffering from, or in danger of contracting, septic shock.

US PAT NO: 5,183,828 [IMAGE AVAILABLE]
DATE ISSUED: Feb. 2, 1993 L1: 3 of 8

Polyhydroxybenzoic acid derivatives Bartholomeus van't Riet, 3419 Noble Ave., Richmond, VA INVENTOR:

23222 **Howard L. Elford**, 3313 Gloucester Rd., Richmond, VA

Galen L. Wampler, 6938 Chamberlayne Rd., Mechanicsville, VA 23111

APPL-NO: 07/555,834 DATE FILED: Jul. 20, 1990

ART-UNIT: 125 PRIM-EXMR: Frederick E. Waddell ASST-EXMR: T. J. Criares LEGAL-REP: James L. Rowe

US PAT NO: 5.183,828 [IMAGE AVAILABLE]

1.1:3 of 8

Polyhydroxy-substituted benz, phenylacet and mandelamidines, amidates, amidoximes and hydroxyamidoximes--ribonucleotide reductase inhibitors, and free radical scavengers.

US PAT NO: 4,942,253 [IMAGE AVAILABLE] DATE ISSUED: Jul. 17, 1990 L1: 4 of 8 Polyhydroxybenzoic acid derivatives INVENTOR: Bartholomeus van't Riet, 3419 Noble Ave., Richmond, VA 23222 **Howard L. Elford**, 3343 Gloucester Rd., Richmond, VA Galen L. Wampler, 6938 Chamberlayne Rd., Mechanicsville, VA 23111 APPL-NO: 06/907,562 DATE FILED: Sep. 15, 1986 ART-UNIT: 122

ART-UNIT: PRIM-EXMR: Anton H. Sutto LEGAL-REP: James L. Rowe

US PAT NO: 4,942,253 [IMAGE AVAILABLE]

L1: 4 of 8

ABSTRACT:

Polyhydroxy-substituted benz, phenylacet and mandelamidines, amidates, amidoximes and hydroxyamidoximes--ribonucleotide reductase inhibitors, and free radical scavengers.

US PAT NO: 4,623,659 [IMAGE AVAILABLE] DATE ISSUED: Nov. 18, 1986 L1: 5 of 8 TITLE: Polyhydroxybenzoic acid derivatives
INVENTOR: Bartholomers Bartholomeus van't Riet, 3419 Noble Ave., Richmond, VA 23222 **Howard L. Elford**, 3313 Gloucester Rd., Richmond, VA Galen L. Wampler, 6938 Chamberlayne Rd., Mechanicsville, VA 23111 APPL-NO: 06/497,370 DATE FILED: May 23, 1983 ART-UNIT: 126

PRIM-EXMR: Natalie Trousof ASST-EXMR: L. Hendriksen LEGAL-REP:

Charles W. Ashbrook, James L. Rowe

US PAT NO: 4.623.659 [IMAGE AVAILABLE]

L1: 5 of 8

Polyhydroxy-substituted benz, phenylacet and mandelamidines, amidates, amidoximes and hydroxyamidoximes--ribonucleotide reductase inhibitors, and free radical scavengers.

US PAT NO: 4,448,730 [IMAGE AVAILABLE] L1: 6 of 8 DATE ISSUED: May 15, 1984

Hydroxybenzohydroxamic acids, benzamides and esters and related compounds as ribonucleotide reductase inhibitors Bartholomeus van't Riet, 3419 Noble Ave., Richmond, VA INVENTOR:

23222

Howard L. Elford, 3313 Gloucester Rd., Richmond, VA 23227

Galen L. Wampler, 6938 Chamberlayne Rd., Mechanicsville, VA 23111

APPL-NO: 06/370,023 DATE FILED: Apr. 20, 1982 ART-UNIT: 126 PRIM-EXMR: Paul J. Killos

James L. Rowe, Arthur R. Whale LEGAL-REP:

US PAT NO: 4,448,730 [IMAGE AVAILABLE] L1: 6 of 8

ABSTRACT:

Di, tri and tetrahydroxybenzohydroxamic acids, amides and the corresponding di, tri and tetrahydroxy substituted phenylalkanohydroxamic acids, amides and phenyl esters, ribonucleotide reductase inhibitors.

US PAT NO: 4,394,389 [IMAGE AVAILABLE] L1: 7 of 8
DATE ISSUED: Jul. 19, 1983
TITLE: Hydroxybenzohydroxamic acids, benzamides and esters as

ribonucleotide reductase inhibitors

INVENTOR: Bartholomeus van't Riet, 3419 Nobel Ave., Richmond, VA 23222 **Howard L. Elford**, 3313 Gloucester Rd., Richmond, VA

Galen L. Wampler, 6938 Chamberlayne Rd., Mechanicsville,

VA 23111

APPL-NO: 06/247,171 DATE FILED: Mar. 24, 1981 ART-UNIT: 117 PRIM-EXMR: Thomas A. Waltz James L. Rowe, Arthur R. Whale LEGAL-REP: US PAT NO: 4.394.389 [IMAGE AVAILABLE] L1: 7 of 8 Di and trihydroxybenzohydroxamic acids, amides, alkyl substituted amides and phenyl esters, ribonucleotide reductase inhibitors. US PAT NO: 4,263,322 [IMAGE AVAILABLE] L1: 8 of 8 DATE ISSUED: Apr. 21, 1981 Hydroxy benzohydroxamic acids and benzamides INVENTOR: Bartholomeus van't Riet, 3419 Noble Ave., Richmond, VA 23222 **Howard L. Elford**, 3313 Gloucester Rd., Richmond, VA 23227 Galen L. Wampler, 6938 Chamberlayne Rd., Mechanicsville, VA 23111 APPL-NO: 06/016,472 DATE FILED: Mar. 1, 1979 ART-UNIT: 117 PRIM-EXMR: Thomas A. Waltz James L. Rowe, Arthur R. Whale LEGAL-REP: US PAT NO: 4,263,322 [IMAGE AVAILABLE] L1: 8 of 8 ABSTRACT: Di or trihydroxybenzohydroxamic acids or N-substituted benzamides, inhibitors or ribonucleotide reductase. = > s ?hydroxybenzo? or (?hydroxy benzo?) TERM 'BENZO?' EXCEEDED TRUNCATION LIMITS - SEARCH ENDED = > s ?hydroxybenzo? or (?hydroxy (w) (benzohydro? or benzoic or benzoate)) 29667 ?HYDROXYBENZO? 151207 ?HYDROXY 273 BENZOHYDRO? 40894 BENZOIC 35208 BENZOATE 3094 ?HYDROXY (W) (BENZOHYDRO? OR BENZOIC OR BENZOATE) 31133 ?HYDROXYBENZO? OR (?HYDROXY (W) (BENZOHYDRO? OR BENZOIC OR BENZOATE)) = > s nf? or (nuclear factor) 15491 NF? 62536 NUCLEAR 268223 FACTOR 205 NUCLEAR FACTOR (NUCLEAR(W)FACTOR) 15563 NF? OR (NUCLEAR FACTOR) = > s l2 (p) l3 19 L2 (P) L3 = > d 14 1-19 bib ab kwic US PAT NO: 5,876,930 [IMAGE AVAILABLE] L4: 1 of 19 DATE ISSUED: Mar. 2, 1999 Hybridization assay using self-quenching fluorescence INVENTOR: Kenneth J. Livak, San Jose, CA

Susan J. A. Flood, Fremont, CA Jeffrey Marmaro, Aurora, CO Khairuzzaman Bashar Mullah, Union, CA Perkin-Elmer Corporation, Foster, CA (U.S. corp.) ASSIGNEE: 08/558,303 APPL-NO: DATE FILED: Nov. 15, 1995 ART-UNIT: PRIM-EXMR: 164 W. Gary Jones ASST-EXMR: Jezia Riley LEGAL-REP: Wilson Sonsini Goodrich & Rosati US PAT NO: 5,876,930 [IMAGE AVAILABLE] L4: 1 of 19

ABSTRACT

A hybridization assay is provided which uses an oligonucleotide probe which includes a fluorescent reporter molecule and a quencher molecule capable of quenching the fluorescence of the reporter molecule. The oligonucleotide probe is constructed such that the probe exists in at least one single-stranded conformation when unhybridized where the

quencher molecule is near enough to the reporter molecule to quench the fluorescence of the reporter molecule. The oligonucleotide probe also exists in at least one conformation when hybridized to a target polynucleotide where the quencher molecule is not positioned close enough to the reporter molecule to quench the fluorescence of the reporter molecule. By adopting these hybridized and unhybridized conformations, the reporter molecule and quencher molecule on the probe exhibits different fluorescence signal intensities when the probe is hybridized and unhybridized As a result, it is possible to determine whether the probe is hybridized or unhybridized based on a change in the fluorescence intensity of the reporter molecule, the quencher molecule, or a combination thereof. In addition, because the probe can be designed such that the quencher molecule quenches the reporter molecule when the probe is not hybridized, the probe can be designed such that the reporter molecule exhibits limited fluorescence until the probe is either hybridized or digested.

DETDESC:

DETD(45)

Compound 2: N,N-Diisopropylethylamine (1.1 g, 1.48 mL, 8.52 mmol), 1-**hydroxybenzotriazol** (420 mg, 3.1 mmol) and (2-(1H-benzotriazol-1yl)-1,1,3,3-tetramethyluronium hexafluorophosphate (1.17 g, 3.1 mmol) were added to a stirred solution of **Nfmoc**-.epsilon.-aminocaproic acid (1 g, 2.84 mmol) in DMF (30 mL) at room temperature. After 15 min DL-homoserine (1.35 g, 11.36 mmol).

US PAT NO: 5,723,591 [IMAGE AVAILABLE]
DATE ISSUED: Mar. 3, 1998
TITLE: Self-quenching fluorescence probe
INVENTOR: Kenneth J. Livak, San Jose, CA L4: 2 of 19 Susan J.A. Flood, Fremont, CA Jeffrey Marmaro, Aurora, CO Khairuzzaman Bashar Mullah, Union City, CA ASSIGNEE: Perkin-Elmer Corporation, Foster City, CA (U.S. corp.) APPL-NO: 08/559.405 DATE FILED: Nov. 15, 1995 ART-UNIT: PRIM-EXMR: Ardin H. Marschel ASST-EXMR: Jezia Riley Wilson Sonsini Goodrich & Rosati LEGAL-REP: US PAT NO: 5,723,591 [IMAGE AVAILABLE] L4: 2 of 19

ABSTRACT:

An oligonucleotide probe is provided which includes a fluorescent reporter molecule and a quencher molecule capable of quenching the fluorescence of the reporter molecule. The oligonucleotide probe is constructed such that the probe exists in at least one single-stranded conformation when unhybridized where the quencher molecule is near enough to the reporter molecule to quench the fluorescence of the reporter molecule. The oligonucleotide probe also exists in at least one conformation when hybridized to a target polynucleotide where the quencher molecule is not positioned close enough to the reporter molecule to quench the fluorescence of the reporter molecule. By adopting these hybridized and unhybridized conformations, the reporter molecule and quencher molecule on the probe exhibit different fluorescence signal intensities when the probe is hybridized and unhybridized. As a result, it is possible to determine whether the probe is hybridized or unhybridized based on a change in the fluorescence intensity of the reporter molecule, the quencher molecule, or a combination thereof. In addition, because the probe can be designed such that the quencher molecule quenches the reporter molecule when the probe is not hybridized, the probe can be designed such that the reporter molecule exhibits limited fluorescence until the probe is either hybridized or digested.

DETDESC:

DETD(48)

Compound 2: N,N-Diisopropylethylamine (1.1 g, 1.48 mL, 8.52 mmol), 1-**hydroxybenzotriazol** (420 mg, 3.1 mmol) and (2-(1H-benzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate (1.17 g, 3.1 mmol) were added to a stirred solution of **Nfmoc**-.epsilon.-aminocaproic acid (1 g, 2.84 mmol) in DMF (30 mL) at room temperature. After 15 min DL-homoserine (1.35 g, 11.36 mmol).

US PAT NO: 5,716,628 [IMAGE AVAILABLE]

DATE ISSUED: Feb. 10, 1998

TITLE: Synergistic biocide composition containing pyrithione plus an additive

INVENTOR: Robert T. Vinopal, Mansfield, CT

John D. Nelson, Jr., Bethlehem, CT

Michael W. Glynn, Darien, CT

Robert W. Coughlin, Storrs, CT

Robert F. Vieth, Manchester, CT

Jon R. Geiger, West Harnford, CT

ASSIGNEE: The University of Connecticut, Storrs, CT (U.S. corp.)

APPL-NO: 08/688.136 DATE FILED: Jul. 29, 1996 ART-UNIT: 124 PRIM-EXMR: Paul J. Killos Dale LynnWiggin & Dana Carlson LEGAL-REP: US PAT NO: 5,716,628 [IMAGE AVAILABLE] L4: 3 of 19 ABSTRACT: Disclosed herein is an antimicrobial composition characterized by synergistic antibacterial and antifungal efficacy and comprising a synetgistic antioacterial and antituding efficiency and comprising a pyrithione salt or pyrithione acid, and at least one compound selected from the group consisting of benzyl and lower alkyl esters of para-hydroxybenzoic acid, salts thereof, carboxylic acids, salts thereof, and combinations thereof. Also disclosed is a method of imparting antimicrobial activity to a composition comprising water or an organic solvent which comprises adding thereto an antimicrobially effective amount of the above-described antimicrobial composition. DETDESC: DETD(5) TABLE 1 Synergistic Antibacterial Effects of Sodium Pyrithione (**NFT**) Mixtures MIC of Mixture (ppm).sup.4 Test Compound NPT, Ratio FIC Test Compound (TC/NPT) ("TC") ppm Index.sup.b. >12/1 <0.53 >6/1 <0.52 781 64 64 391 methyl ester of p-2500 0 **hydroxybenzoate** **hydroxyound methyl ester of p-425 I6 39/1 0.38 **hydroxybenzoate** methyl ester of p-313 32 313 10/1 0.38 **hydroxyound methyl ester of p-156 32 5/1 0.31 **hydroxybenzoate** methyl ester of p-2/1 0.28 **hydroxybenzoate** methyl ester of p-1/2 0.52 **hydroxybenzoate** methyl ester of p-64 1/3 0.51 20 **hydroxybenzoate** methyl ester of p-1/6 0.50 10 **hydroxybenzoate** none 0 64 sorbic acid 4096 2048 16 128/1 0.75 1024 32. . . US PAT NO: 5,688,828 [IMAGE AVAILABLE] L4: 4 of IIDATE ISSUED: Nov. 18, 1997
TITLE: Use of N,N'-bis(mercaptoacetyl) hydrazine derivatives as L4: 4 of 19 anticataract agents
R: Mark R. Hellberg, Arlington, TX INVENTOR: William H. Garner, Southlake, TX Jaime E. Dickerson, Jr., Fort Worth, TX Marjorie F. Lou, Lincoln, NE E: Alcon Laboratories, Inc., Fort Worth, TX (U.S. corp.) ASSIGNEE: APPL-NO: 08/690,610 DATE FILED: Jul. 31, 1996 ART-UNIT: 125 PRIM-EXMR: Zohreh Fay LEGAL-REP: Michael C. Mayo US PAT NO: 5,688,828 [IMAGE AVAILABLE] L4: 4 of 19

ABSTRACT:

Compositions containing certain sulfur containing compounds and methods of use in the treatment and prevention of cataracts is disclosed.

DETDESC:

DETD(47)

%) Purpose

Compound 0.1 active ingredient Sodium chloride, USP Boric acid, USP 0.4 preservative Methyl p-**hydroxybenzoate** preservative 0.002 USP Chlorobutanol, USP 0.03 Sodium hydroxide, **NF** q.s. pH adjustment Hydrochloric acid, **NF** pH adjustment q.s. Water for injection, USP vehicle q.s.

US PAT NO: 5,688,529 [IMAGE AVAILABLE]
DATE ISSUED: Nov. 18, 1997 TITLE: Mycophenolate mofetil high dose oral suspensions INVENTOR: Deborah Marilyn Lidgate, Los Altos, CA Li-hua Wang-Kessler, Palo Alto, CA Bindu Joshi, Milpitas, CA Sayee Gojanan Hegde, Los Angeles, CA Leo Gu, Saratoga, CA E: Syntex (U.S.A) Inc., Palo Alto, CA (U.S. corp.) ASSIGNEE: APPL-NO: 08/412,645 DATE FILED: Mar. 29, 1995 ART-UNIT: PRIM-EXMR: Thurman K. Page ASST-EXMR: James M. Spear Heller Ehrman White & McAuliffe LEGAL-REP: US PAT NO: 5,688,529 [IMAGE AVAILABLE] L4: 5 of 19 ABSTRACT:

High dose, dry granulations or powder blends and aqueous oral suspensions of mycophenolate mofetil or mycophenolic acid, contain: active compound (7.5-30%), suspending/viscosity agent, sweetener, flavor, buffer (to a pH of 5-7.5), and optionally contain flavor enhancer, wetting agent, antimicrobial agent and color.

SUMMARY:

BSUM(40)

Antimicrobial agents useful in the formulations of the invention include, for example: sodium benzoate; sodium methyl paraben (preferably **NF**: sodium methyl paraben); methyl paraben (preferably **NF**: methyl paraben, or BP: methyl **hydroxybenzoate**, or EP: methylis **parahydroxybenzoas**); propylparaben (preferably **NF**: propylparaben, or BP/EP: propyl **hydroxybenzoate**); and potassium sorbate (preferably **NF** or BP).

US PAT NO: 5,686,450 [IMAGE AVAILABLE] L4: 6 of 1
DATE ISSUED: Nov. 11, 1997
TITLE: Use of N,N'-bis(mercaptoacetyl) hydrazine derivatives as L4: 6 of 19 INVENTOR: Mark R. Hellberg, Arlington, TX
William H. Garner, Southlake, TX
Jaime E. Dickerson, Jr., Fort Worth, TX Marjorie F. Lou, Lincoln, NE ASSIGNEE: Alcon Laboratories, Inc., Fort Worth, TX (U.S. corp.) 08/472,452 APPL-NO: DATE FILED: Jun. 7, 1995 ART-UNIT: 125 PRIM-EXMR: Zohreh Fay Michael C. Mayo LEGAL-REP: US PAT NO: 5,686,450 [IMAGE AVAILABLE] L4: 6 of 19

ABSTRACT:

Compositions containing certain sulfur containing compounds and methods of use in the treatment and prevention of cataracts is disclosed.

DETDESC:

DETD(38)

%) Purpose

Compound 0.1 active ingredient
Sodium chloride, USP
0.7 tonicity
Boric acid, USP

0.4 preservative Methyl p-**hydroxybenzoate** 0.002 preservative Chlorobutanol, USP 0.03 Preservative Sodium hydroxide, **NF** pH adjustment q.s. Hydrochloric acid. pH adjustment q.s. Water for injection, USP q.s. vehicle

US PAT NO: 5,357,636 [IMAGE AVAILABLE]

L4: 7 of 19

DATE ISSUED: Oct. 25, 1994

TITLE: Flexible protective medical gloves and methods for their

INVENTOR: Karl P. Dresdner, Jr., 235 W. 48th St., Apt. #18N, New

York City, NY 10036 Kenneth H. Dangman, 400 Riverside Dr., Apt. #1A, New York City, NY 10032

Edward A. Jazlowiecki, 15 Sachems Trail, West Simsbury, CT 06092

APPL-NO: 07/906,829

DATE FILED: Jun. 30, 1992

ART-UNIT: 247

Clifford D. Crowder PRIM.FYMR-

ASST-EXMR: Amy B. Vanatta

US PAT NO: 5,357,636 [IMAGE AVAILABLE]

1.4: 7 of 19

A flexible protective medical glove containing a non-liquid antiseptic composition and methods for its use are disclosed. The glove comprises a thin inner layer and a thin outer layer of material; preferably the outer layer is a more elastic and less plastic layer than the inner layer. A compartment between the layers of the glove is capable of providing a non-liquid antiseptic composition which comprises an antiseptic in a non-liquid composition. The non-liquid antiseptic composition may also contain a surface-active agent, an algesic agent, a colorant, a vasoconstrictive agent, an odorant, or a viscosity-modifying agent. An object puncturing the glove wall can become coated with the non-liquid antiseptic composition and can automatically transfer some of the antiseptic composition from the glove onto the hand and into a hand wound should the object cause a wound; useful as an immediate preventative antiseptic treatment to help to decontaminate the hand and hand wound of infectious pathogens that may have been transferred there by the object. The treatment can help to protect a gloved individual such as a surgeon, a medical doctor, a health care worker, a law enforcement officer, a dentist or any worker whose work may place them at some risk of becoming contaminated through the hands by an infectious pathogen including the AIDS virus or hepatitis B virus.

DETDESC:

DETD(39)

chlorhexidine acetate, chlorhexidine hydrochloride, chlorhexidine, other chlorhexidine salts, other hexamethylenebis biguanides, octoxynol, nonoxynol-9, methanol, ethanol, isopropanol, allyl alcohol, rubbing alcohol **NF**, sodium hypochlorite, potassium hypochlorite, calcium hypochlorite, magnesium hypochlorite, sodium dichloroisocyanurate, sodium perborate **NF**, sodium hydroxide, potassium hydroxide, magnesium hydroxide, calcium hydroxide, ammonia, ammonium hydroxide, lithium hydroxide, barium hydroxide, silver hydroxide, other metal hydroxides, . . . eucalyptus oil, glycobiarsol, gramicidin, hexyl resorcinol, methylene blue, peppermint oil, phenylethyl alcohol, phenyl salicylate, methyl salicylate, pine tar, pine oil **NF**, pine oil emulsion, tertiary terpene alcohols, secondary terpene alcohols, alpha-terpineol, borneol, fenchyl alchol, o-methylchavicol, polymixin B sulfate, colistin, chloramphenicol, tetracycline,. . . sulfisoxazole diolamine, sulfacetamide sodium, gentamycin sulfate, amphotericin B, tobramycin, a penicillin, a cephalosporin, salicylic acid, rrichloroacetic acid, benzoic acid, pyrogallol **NF** X, pyrogallic acid, sodium benzoate, boric acid, sodium borate, lactic acid, sodium lactate, chloramine, chloramine T, silver nitrate, ammoniacal silver. mercuric iodide red, mercuric oxide red, strontium iodide, lithium iodide, magnesium iodide, zinc iodide, silver iodide, selenium iodide, thymol iodide **NF** X, dithymol diiodide, iodinated derivatives of thymol, other iodide salts, povidone-iodine, iodoform, iodinated organ compounds, iodol, iodopyrrol, other iodophors, chlorinated lime, bromide salts, sodium bromide, merbromin **NF**, other bromophors, other brominated chemicals, sodium fluoride and other fluorinated chemicals and fluorophors, Lysol, Nonidet P40, phenyl mercuric acetate, potassium mercuric iodide, proflavine hemisulfate, 3,6-diaminoacridine bisulfate, formaldehyde, glutaraldehyde, parsformaldehyde, butyl **hydroxybenzoate**, mercurous chloride, iodochlorhydroxyquin, zinc nitrate, zinc sulfate, cadmium sulfate, thimerosal **NF**, zinc oxide,

zinc acetate, zinc chloride, silver nitrate, silver sulfadiazine, hydrogen peroxide, urea hydrogen peroxide, hydrogen peroxide carbamide, benzoyl peroxide, . . . perchlorite, sodium perchlorite, calcium perchlorite, magnesium perchlorite, zinc perchlorite, zinc peroxide, zinc carbonate, zinc hydroxide, zinc sulfate, succinyl peroxide, succinchlorimide **NF** IX, N-Chloro-succinimide, potassium permaganate, sodium chlorate, potassium chlorate, phenol, sodium phenolate, domiphen bromide, salicylic acid, bismuth-formic-iodide, bismuth subgallate, bacitracin zinc,. . . hydroxynalidixic acid, pipemidic acid, norfloxacin, norfloxacin hydrochloride, other quinolones, 8-hydroxyquinoline sulfate, sodium phenolate, thyme oil, o-cresol, m-cresol, metacresylacetate, p-cresol, cresol **NF**, 4-chloro-m-cresol, 4-chloro-S,5-xylenol, saponified cresol solution **NF**, methylphenol, ethyl phenol, other alkyl phenols, o-phenyl phenol, other aryl phenols, bis-phenols, phenyl-mecuric chloride, phenylmecuric borate, resorcinol, resorcinol monoactetate **NF**, orthophenylphenol, chloroxylenol, hexyl-resorcinol, parachlorophenol, paratertiary-amylphenol, thymol, chlorothymol **NF**, menthol, butylparaban, ethylparaben, methylparaben, propylparaben, triclosan, bithionol **NF**, o-benzyl-p-chlorophenol, hexachlorophene, poloxamer 188, benzalkonium chloride where the alkyl groups attached to the nitrogen represent any alkyl from CH.sub.3 to. C.sub.18 H.sub.37, methylbenzethonium chloride, cetrimonium bromide, abikoviromycin, acetylenedicarboxamide, acetyl sulfamethoxypyrazine, triclobisonium chloride, undecoylium chlorideiodine, coal tar solution, furazolidone, nifuroxime, nitrofurazone **NF**, nitromersol **NF**, oxychlorosene, sodium oxychlorosene, parachlorophenol **NF**, camphorated parachlorophenol **NF**, phenylmercuric nitrate **NF**, gentian violet USP, hexamethylpara-rosaniline chloride, rosaniline chloride pentamethylpararosaniline chloride, methylrosaniline chloride, tetramethylpararosaniline chloride, nonylphenoxypolyethoxyethanol, methoxypolyoxyetheneglycol 550 laurate, oxyquinoline benzoate, p-triisopropylphenoxypolyethoxy-ethanol, halazone **NF**, dichloramine-T, benzethonium chloride, econazole, cetylpyridinium chloride, methylbenzethonium chloride, cetyldimethylbenzylammonium chloride, dichlorobenzalkonium chloride, domiphen bromide, triclocarban, clotrimazole, ciclopirox olamine, undecylenic acid, . . acid acriflavine, 5-aminoacridine hydrochloride monohydrate, malachite green G, dodecyltrimethylammonium bromide, tetradecyltrimethylammonium bromide, dequalinium chloride BP, dibromopropamidine isethionite, hexadecyltrimethylammmonium bromide, chloroazodin **NF** X, N-chloro-p-toluenesulfonamidosodium, 4-[(dichloroamino)sulfonyl]-benzoic acid, methenamine, methenamine mandelate, methenamine hippurate, octoxynol 9, phenazopyridine hydrochloride, 9-aminoacridine hydrochloride, bismuth tribromophenate, p-tert-butylphenol, cetyldimethylethylammonium bromide, . .

CLAIMS:

CLMS(5)

5. . . from the group consisting of chlorhexidine gluconate, chlorhexidine acetate, chlorhexidine hydrochloride, octoxynol nonoxynol-9, methanol, ethanol, isopropanol, allyl alcohol, rubbing alcohol **NF**, sodium hypochlorite, potassium hypochlorite, calcium hypochlorite, magnesium hypochlorite, sodium dichloroisocyanurate, sodium perborate **NF**, sodium hydroxide, potassium hydroxide, magnesium hydroxide, calcium hydroxide, ammonia, ammonium hydroxide, lithium hydroxide, barium hydroxide, silver hydroxide, sodium tetradecyl sulfate, . . . eucalyptus oil, glycobiarsol, gramicidin, hexyl resorcinol, methylene blue, peppermint oil, phenylethyl alcohol, phenyl salicylate, methyl salicylate, pine tar, pine oil **NF**, alpha-terpineol, borneol, fenchyl alcohol, o-methylchavicol, polymixin B sulfate, salicylic acid, trichloroacetic acid, benzoic acid, pyrogallol **NF** X, pyrogallic acid, sodium benzoate, boric acid, sodium borate, lactic acid, sodium lactate, ohiofamine, chloramine T, silver nitrate, ammoniacal silver. . . mecuric iodide red, mecuric oxide red, strontium iodide, lithium iodide, magnesium iodide, zinc iodide, silver iodide, selenium iodide, thymol iodide **NF** X, dithymol diiodide, povidone-iodine, iodoform, iodol, iodopyrrol, chlorinated lime, potassium bromide, sodium bromide, merbromin **NF**, sodium fluoride, potassium fluoride, phenyl mercuric acetate, potassium mecuric iodide, proflavine hemisulfate, 3,6-diaminoacridine bisulfate, formaldehyde, glutaraldehyde, paraformaldehyde, butyl **hydroxybenzoate**, mercurous chloride, iodochlorhydroxyquin, zinc nitrate, zinc sulfate, cadmium sulfate, thimerosal **NF**, zinc oxide, zinc acetate, zinc chloride, silver nitrate, silver sulfadiazine, hydrogen peroxide, urea hydrogen peroxide, hydrogen peroxide carbamide, benzoyl peroxide,. . . perchlorite, sodium perchlorite, calcium perchlorite, magnesium perchlorite, zinc perchlorite, zinc peroxide, zinc carbonate, zinc hydroxide, zinc sulfate, succinyl peroxide, succinchlorimide **NF** IX, N-Chlorosuccinimide, potassium permanganate, sodium chlorate, potassium chlorate, phenol, camphorated phenol, phenol glycerin, chloroxylenol, 4-chloro-3,5-xylenol, sodium phenolate, domiphen bromide, salicylic. . . zinc sulfocarbolate, hydroxynalidixic acid, pipemidic acid, norfloxacin, norfloxacin hydrochloride, 8-hydroxyquinoline sulfate, sodium phenolate, thyme oil, o-cresol, m-cresol, metacresylacetate, p-cresol, cresol **NF**, 4-chloro-m-cresol, 4-chloro-3,5-xylenol, saponified cresol solution **NF**, methylphenol, ethyl phenol, other alkyl phenols, o-phenyl phenol, other aryl phenols, bisphenols, phenyl-mecuric chloride, phenylmecuric borate, resorcinol, resorcinol monoactetate **NF**,

orthophenylphenol, chloroxylenol, hexylresorcinol, parachlorophenol, paratertiary-amylphenol, thymol, chlorothymol **NF**, butylparaba ethylparaben, methylparaben, propylparaben, triclosan, bithionol **NF** o-benzyl-p-chlorophenol, hexachlorophene, poloxamer 188, a benzalkonium chloride wherein the alkyl groups attached to the nitrogen represent an alkyl from CH.sub.3 to C.sub.18 H.sub.37, triclobisonium chloride, undecoylium chlorideiodine, coal tar solution, furazolidone, nifuroxime, nitrofurazone **NF**, nitromersol **NF**, oxychlorosene, sodium oxychlorosene, parachlorophenol **NF**, camphorated parachlorophe **NF**, phenylmercuric nitrate **NF**, gentian violet USP, hexamethylpara-rosaniline chloride, rosaniline chloride, pentamethylpararosaniline chloride, methylrosaniline chloride, tetramethylpararosaniline chloride, nonylphenoxypolyethoxyethanol, methoxypolyoxyetheneglycol 550 laurate, oxyquinoline benzoate, p-triisopropylphenoxypolyethoxy-ethanol, halazone "*NF**, dichloramine-T, benzethonium chloride, econazole, cetylpyridinium chloride, methylbenzethonium chloride, cetyldimethylbenzylammontum chloride, dichlorobenzalkonium chloride, domiphen bromide, triclocarban, clotrimazole, ciclopirox olamine, undecylenic acid, . . . acid acr]flavine, 5-aminoacridine hydrochloride monohydrate, malachite green G, dodecyltrimethylammonium bromide, tetradecyltrimethyl-ammonium bromide, dequalinium chloride BP, dibromopropamidine isethionite, hexadecyltrimethylammmonium bromide, chloroazodin **NF** X, N-chloro-p-toluenesulfonamidosodium, 4-[(dichloroamino)sulfonyl]-benzoic acid, methenamine, methenamine mandelate, methenamine hippurate, octoxynol 9, phenazopyridine hydrochloride, 9-aminoacridine hydrochloride, bismuth tribromophenate, p-tert-butylphenol, cetyldimethylethylammonium bromide,. . .

```
US PAT NO: 5,082,651 [IMAGE AVAILABLE]
                                                              L4: 8 of 19
DATE ISSUED: Jan. 21, 1992
             Pharmaceutical compositions
INVENTOR:
          OR: John N. C. Healey, Hitchin, England
Marshall Whiteman, Baldock, England
ASSIGNEE:
               Smith Kline & French Laboratories Limited, Welwyn Garden
           City, England (foreign corp.)
07/514,634
APPL-NO:
DATE FILED: Apr. 25, 1990
ART-UNIT:
                125
                 Shep K. Rose
PRIM-EXMR:
LEGAL-REP:
                 Linda E. Hall, Stuart R. Suter, Edward T. Lentz
               5,082,651 [IMAGE AVAILABLE]
                                                              1.4: 8 of 19
US PAT NO:
Pharmaceutical compositions suitable for intra-rectal administration in
the form of a foam are described which comprise a therapeutically effective amount of 5-aminosalicylic acid, a pharmaceutically acceptable
aqueous carrier therefor, and means for generating a foam.
DETDESC:
DETD(2)
PROPELLANT)
                     Weight
5-Aminosalicylic Acid 15.0 150.0
Polysorbate 80
                     0.25 2.5
Emulsifying Wax ('Polawax **NF**')
                 0.5
                        5.0
Colloidal Silicon Dioxide (`Aerosil 200`)
0.5 5.0
Sodium Metabisulphite 0.3 3.0
Disodium Edetate, dihydrate
                 0.1
**Methylparahydroxybenzoate**
                        2.0
                 0.2
 **Propylparahydroxybenzoate**
                 0.04 0.4
Disodium Hydrogen Orthophosphate, 12H.sub.2 O
1.19 11.9
Sodium Dihydrogen Orthophosphate, 2H.sub.2 O
                 0.52
Glycerol. . .
DETDESC:
DETD(3)
                % (w/w)
                     (g)
5-Aminosalicylic Acid 25.0 150.0
Sorbitan mono-oleate (`Span 80`)
0.25 1.5
Emulsifying Wax ('Polawax **NF**')
```

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Colloidal Silicon Dioxide ('Aerosil 200')
               0.5
                      3.0
Sodium Metabisulphite 0.3
Disodium Edetate, dihydrate
               0.1
                      0.6
**Methylparahydroxybenzoate*
               0.2
**Propylparahydroxybenzoate
                      0.24
               0.04
Disodium Hydrogen Orthophosphate, 12H.sub.2 O
               1.19
                     7.14
Sodium Dihydrogen Orthophosphate, 2H.sub.2 O
               0.52 3.12
Glycerol. . .
DETDESC:
DETD(4)
CONCENTRATE
(without Addition of Propellant)
                   Weight
              % (w/w)
                   (g)
5-Aminosalicylic Acid 25.0
                             250.0
Emulsifying Wax ('Polawax *
               0.75
                      7.5
Sodium Metabisulphite 0.3
Disodium Edetate, dihydrate
               0.1
**Methylparahydroxybenzoate*
               0.2
**Propylparahydroxybenzoate**
               0.04 0.4
Disodium Hydrogen Orthophosphate, 12H.sub.2 O
               1.19 11.9
Sodium Dihydrogen Orthophosphate, 2H.sub.2 O
0.52 5.2
Water. . .
DETDESC:
DETD(6)
FOAM
                   Weight
5-Aminosalicylic Acid 25.0 250.0
Polysorbate 80 0.25 2.50
Emulsifying Wax ('Polawax **NF**')
               0.5 5.0
Colloidal Silicon Dioxide ('Aerosil 200')
               0.5
                    5.0
Sodium Metabisulphite 0.3
Disodium Edetate, dihydrate
**Methylparahydroxybenzoate**
               0.2
                      2.0
**Propylparahydroxybenzoate**
               0.04 0.4
Disodium Hydrogen Orthophosphate, 12H.sub.2 O
               1.19 11.9
Sodium Dihydrogen Orthophosphate, 2H.sub.2 O
               0.52
Glycerol. . .
DETDESC:
DETD(7)
   Weight
              (w/w) (g)
5-Aminosalicylic Acid 22.50 2.475
Polysorbate 20 (Tween 20)
               10.00 1.100
Emulsifying Wax ('Polawax **NF**')
0.40 0.044
Colloidal Silicon Dioxide (Aerosil 200)
0.40 0.044
Sodium Metabisulphite 0.30 0.033
Disodium Edetate, dihydrate
               0.10 0.011
**Methylparahydroxybenzoate**
               0.20 0.022
**Propylparahydroxybenzoate**
```

0.04 0.0044

Disodium Hydrogen Orthophosphate, 12H.sub.2 O 1.19 0.131

Sodium Dihydrogen Orthophosphate, 2H.sub.2 O

0.52 0.057

Glycerol. . .

US PAT NO: 5,041,280 [IMAGE AVAILABLE]

DATE ISSUED: Aug. 20, 1991
TITLE: Toothpaste composition for stain removal

L4: 9 of 19

INVENTOR: Irwin E. Smigel, New York, NY Epilady USA, Inc., Culver City, CA (U.S. corp.) ASSIGNEE:

APPL-NO: 07/103.533

DATE FILED: Oct. 1, 1987 ART-UNIT: 189

PRIM-EXMR: F. T. Moezie LEGAL-REP:

Darby & Darby

US PAT NO: 5,041,280 [IMAGE AVAILABLE] L4: 9 of 19

ABSTRACT:

A toothpaste composition having the following ingredients, by weight:

Dicalcium Phosphate dihydrate:

From 1.0% to 50%

Calcium Carbonate From 1.0% to 50%

Sodium Bicarbonate From 1.0% to 50% Magnesium Carbonate From 1.0% to 25%

Sorbitol 70% Corn Starch

From 1.0% to 50% From 0.5% to 10%

Celtulose Gum From 0.5% to 5.0%
Calcium Peroxide From 0.5% to 5.0%
Lathanol LAL (Sodium Laury)

From 0.1% to 5%

Sulfoacetate)

Aluminum Hydroxide From 0.01 to 1%

Saccharinate (Sodium Salt)

From 0.05% to 2%

Flavoring material From 0.05% to 2%

Alkylparaben From 0.05% to 1.0%

Sodium monofluorophosphate From 0.70% to 0.80%

Titanium Dioxide From 0.1% to 10%

Deionized Water From 10% to 50%

DETDESC:

DETD(7)

0.7

Aluminum Hydroxide

Saccharinate (Sodium Salt) 0.5

Flavoring Material

Consisting of:

Menthol crystals 20%

Oil of Spearmint **NF**

20%

Terpeneless Spearmint 30%

Oil of Peppermint 20%

Oil of Anise (imitation)

10%

Methylparaben 0.5

(Hydrobenzoic acid methyl ester) Propylparaben 0.03

(**Hydroxybenzoic** acid propylester)

Sodium Monofluorophosphate

0.76

Titanium Dioxide 1.0

Deionized Water (to make up 100%)

47.46

US PAT NO: 4,898,728 [IMAGE AVAILABLE]

L4: 10 of 19

DATE ISSUED: Feb. 6, 1990

Process for production of composition containing lecithin and polyvinylpyrrolidone TITLE:

Robert R. Vartan, Bristol, TN
Beecham Group p.l.c., England (foreign corp.) INVENTOR: ASSIGNEE:

APPL-NO: 07/178,487 DATE FILED: Apr. 7, 1988

ART-UNIT: 155

PRIM-EXMR: Joseph L. Schofer ASST-EXMR: Carmen B. Pili-Curtis

LEGAL-REP: Jacobs & Jacobs US PAT NO: 4.898.728 IIMAGE AVAILABLE1

I.4: 10 of 19

A process for the production of a sterile composition containing lecithin and polyvinyl pyrrolidone is disclosed in which the lecithin and polyvinyl pyrrolidone, optionally with one or more preservative powders are admixed in a solvent comprising about 75% to 90% methyl isobutyl ketone and about 10% to 25% isopropyl alcohol, and the solution passed through a millipore filter in order to render the ingredients sterile. The process is usefully employed in the production of injectable compositions of amoxycillin and ampicillin.

SUMMARY:

BSUM(11)

Preferably . . . are also dissolved in the MIBK/IPA solvent system in processes of the invention. A preferred preservative is an ester of p-*hydroxybenzoic** acid, and in particular a mixture of methyl p-hydroxybnnzoate and propyl p-**hydroxybenzoate**. A commercial source of these preservative powders are known as methylparaben **NF** and propylparaben **NF** respectively.

L4: 11 of 19

US PAT NO: 4,883,805 [IMAGE AVAILABLE]
DATE ISSUED: Nov. 28, 1989
TITLE: Stable, Injectable solutions of vinca dimer salts
INVENTOR: Rodney Kasan, Raanana, Israel

Haim Yellin, Ramat-Gan, Israel

Michael Seiffe, Raanana, Israel

Teva Pharmaceutical Industries Ltd., Israel (foreign ASSIGNEE: APPL-NO:

07/078.805

DATE FILED: Jul. 28, 1987

ART-UNIT: 118

PRIM-EXMR: William R. Dixon, Jr.

ASST-EXMR: James M. Hunter Steinberg & Raskin

LEGAL-REP:

US PAT NO: 4,883,805 [IMAGE AVAILABLE]

L4: 11 of 19

ABSTRACT:

A stable, injectable pharmaceutical composition of vinca dimer salts. The compositions are in the form of an aqueous solution comprising per 1 ml of solution:

from about 0.2 to about 2 mg of one or more pharmaceutically acceptable vinca dimer salts;

from about 0.1 to about 1.0 mg of a pharmaceutically acceptable ethylenediamine-tetraacetic acid (EDTA) salt;

acetate buffer in an amount necessary to maintain said aqueous solution at a pH of from about 3.0 to about 5.5; and

from about 1.5 to about 2.5 mg of a preservative selected from methyl paraben, propyl paraben and mixtures thereof.

DETDESC:

DETD(28)

Materials

Vincristine sulfate BP/USP

Plantex, Israel Methyl **hydroxybenzoate** BP/**NF**

Machteshim, Israel

(methyl paraben)

Propyl **hydroxybenzoate** BP/**NF**

Machteshim, Israel (propyl paraben)

Edetate disodium BP/USP

Merck, Germany Sodium Acetate 3H sub 2 O BP/USP

Merck, Germany
Acetic acid BP/**NF** Merck, Germany

L4: 12 of 19

US PAT NO: 4,777,050 [IMAGE AVAILABLE] DATE ISSUED: Oct. 11, 1988

TITLE: Controlled-release dosage form comprising acetaminophen,

pseudoephedrine and dexbrompheniramine INVENTOR: Winston A. Vadino, Whitehouse Station, NJ

ASSIGNEE: Schering Corporation, Kenilworth, NJ (U.S. corp.) 07/029,032 APPL-NO: DATE FILED: Mar. 23, 1987

ART-UNIT: 139

PRIM-EXMR: Michael R. Lusignan

LEGAL-REP: Anita W. Magatti, Stephen I. Miller, James R. Nelson

L4: 13 of 19

The invention relates to a controlled release dosage form comprising three actives: acetaminophen, pseudoephedrine and dexbrompheniramine.

DETDESC:

DETD(8)

Approximate Ingredients g/Batch mg/tablet Hydroxypropyl Methylcellulose 1,440 2910 or 2906 USP Polyethylene glycol 3350 **NF** Methyl p-**hydroxybenzoate** **NF** 14.4 0.12 Propyl p-**hydroxybenzoate** **NF** 10.8 0.09 Purified Water USP (evaporates) (1) Coloring Agent (2)

(1) Sufficient amounts of. . .

US PAT NO: 4,690,776 [IMAGE AVAILABLE] L4:
DATE ISSUED: Sep. 1, 1987
TITLE: Method of manufacture of a toothpaste composition

INVENTOR:

Irwin E. Smigel, 635 Madison Ave., New York, NY 10022

APPL-NO: 06/817,043 DATE FILED: Jan. 8, 1986

ART-UNIT: 223

PRIM-EXMR: Richard D. Lovering

Roberts, Spiecens & Cohen LEGAL-REP:

US PAT NO: 4,690,776 [IMAGE AVAILABLE] L4: 13 of 19

ABSTRACT:

A method of preparing a toothpaste composition by the steps of adding calcium phosphate in an amount of 0.5 to 5% by weight of the composition and sodium perborate in an amount of 0.5 to 5% by weight of the composition to hot water to form a first mixture and agitating the first mixture, adding to the first mixture sorbitol in an amount of 1 to 50% by weight of the composition, cornstarch in an amount of 0.5 to 10% by weight of the composition and aluminum hydroxide in an amount of 0.01 to 1% by weight of the composition to form a second mixture and agitating the second mixture, adding to the second mixture dicalcium phosphate in an amount of 1 to 50% by weight of the composition and sodium monofluoride phosphate in an amount of 0.7 to 0.8% by weight of the composition to form a third mixture and agitating the third mixture, adding to the third mixture sodium bicarbonate, in an amount of 1 to 50% by weight of the composition, gradually adding increments of a flavoring material in an amount of 0.05 to 2% to control any foaming and to facilitate release of gases to form a fourth mixture, and adding to the fourth mixture sodium lauryl sulfoacetate in an amount of 0.1 to 5% and gum in an amount of 0.5 to 5% and agitating the fourth mixture until a homogeneous paste composition is obtained.

SUMMARY:

BSHM(36)

Aluminum Hydroxide Saccharinate (Sodium Salt)

0.5

Flavoring Material

Consisting of:

Menthol crystals 20%

0.1 of Spearment **NF** 20%

Terpeneless Spearmint

30%

Oil of Peppermint 20%

Oil of Anise (imitation)

10%

Methylparaben 0.5

(**Hydroxybenzoic** acid methyl ester) 0.03

Propylparaben

(**Hydroxybenzoic** acid propylester)

Sodium Monofluoride Phosphate

0.76 Titanium Dioxide 1.0

Deionized Water (to make up 100%)

US PAT NO: 4,603,045 [IMAGE AVAILABLE]

L4: 14 of 19

DATE ISSUED: Jul. 29, 1986

Toothpaste for bonded (composite filling material) as well TITLE:

as natural teeth

Irwin E. Smigel, 635 Madison Ave., New York, NY 10022 INVENTOR:

APPL-NO: 06/706,001 DATE FILED: Feb. 27, 1985 ART-UNIT: 123

PRIM-EXMR: Shep K. Rose

LEGAL-REP: Roberts, Spiecens & Cohen

US PAT NO: 4,603,045 [IMAGE AVAILABLE]

L4: 14 of 19

ABSTRACT:

A toothpaste composition consisting essentially of, in percent by weight:

Dicalcium Phosphate, dihydrous: From 1.0% to 50% Calcium Carbonate: From 1.0% to 50%

Sodium Bicarbonate: From 1.0% to 50% Magnesium Carbonate: From 1.0% to 25% Sorbitol 70%: From 1.0% to 50%

Corn Starch: From 0.5% to 10% Cellulose Gum: From 0.5% to 5.0% Calcium Peroxide: From 0.5% to 5% Sodium Perborate: From 0.5% to 5%

Lathanol LAL (Sodium Lauryl Sulfoacetate): From 0.1% to 5% Aluminum Hydroxide: From 0.01 to 1%

Saccharinate (Sodium Salt): From 0.05% to 2% Flavoring material: From 0.05% to 2% Alkylparaben: From 0.05% to 1.0% Sodium Monofluoride Phosphate: From 0.70% to 0.80%

Titanium Dioxide: From 0.1% to 10% Deinonized Water: From 10% to 50%.

SUMMARY:

BSUM(55)

0.7

Aluminum Hydroxide

Saccharinate (Sodium Salt)

0.5

Flavoring Material Consisting of: Menthol crystals 20%
0.1 of Spearment **NF**
Terpeneless Spearmint 30%

Oil of Peppermint 20% Oil of Anise (imitation)

10% Methylparaben 0.5

(Hydroybenzoic acid methyl ester)

Propylparaben 0.03 (**Hydroxybenzoic** acid propylester)

Sodium Monofluoride Phosphate

0.76

Titanium Dioxide 1.0

Deinonized Water (to make up 100%)

47.46

US PAT NO: 4,323,694 [IMAGE AVAILABLE]

L4: 15 of 19

DATE ISSUED: Apr. 6, 1982 TITLE: Benzoic acid esters

INVENTOR: Thomas L. Scala, Jr., West Milford, NJ

ASSIGNEE: Finetex, Inc., Elmwood Park, NJ (U.S. corp.)

APPL-NO: 06/258,801 DATE FILED: Apr. 29, 1981

126 ART-UNIT:

PRIM-EXMR: Paul J. Killos Weingram & Klauber LEGAL-REP:

US PAT NO: 4,323,694 [IMAGE AVAILABLE]

L4: 15 of 19

ABSTRACT:

The benzoic acid ester of an alcohol which is selected from the group consisting of (A) at least one C.sub.2n branched primary alcohol, wherein n is 5 through 9; (B) at least one C.sub.2m+1 branched or linear primary alcohol, wherein m is 4 through 9; and (C) mixtures comprising at least 40% and preferably at least 60% by weight of the members of (A) and (B), with one or more linear primary alcohols of even carbon number chain length. The benzoic acid esters are useful in skin care compositions, e.g., hand cleaners, bath oils, suntan oils, anti-perspirants, perfumes, colognes, cold creams, electric pre-shaves, eye and throat oils, topical pharmaceutical ointments, lipsticks, stick rouges, lotions, skin moisturizers, cleansing creams or after bath splashes or lotions.

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DETDESC:
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DETD(261)

% by wt.

A. Beeswax, white 11.00 Cetyl alcohol 2.50 Cetyl palmitate 2.20
Mineral oil, light, **NF** 28.00 20.60 Benzoate Cerasynt Q (Glyceryl stearate, self-emulsifying).sup.1 0.75 Propyl paraben (propyl p-**hydroxybenzoate**)
0.05 B. Water, purified 32.83

Borax (sodium borate) 0.75 Methyl paraben (methyl p-**hydroxybenzoate**) 0.15

C. Water, purified 1.00 Dowicil 200 (Quaternium-15).sup.2 0.10 D. Fragrance 0.07

.sup.1 Van Dyk. . .

US PAT NO: 4,323,693 [IMAGE AVAILABLE] DATE ISSUED: Apr. 6, 1982 L4: 16 of 19

TITLE: INVENTOR: Benzoic acid ester

Thomas L. Scala, Jr., West Milford, NJ Finetex, Inc., Elmwood Park, NJ (U.S. corp.) 06/257,977 ASSIGNEE:

APPL-NO: DATE FILED: Apr. 27, 1981 ART-UNIT: PRIM-EXMR: 126 Paul J. Killos

LEGAL-REP: Weingram & Klauber

US PAT NO: 4,323,693 [IMAGE AVAILABLE] L4: 16 of 19

A substantially pure benzoic acid ester of isostearyl (C.sub.18) alcohol. The composition of this invention has unique properties in that it is one-greasy, has a low cloud point and pour point, is practically odorless, has low toxicity, and is stable. These properties make such composition useful as a vehicle or carrier, emollient or solubilizer for toiletry and cosmetic formulations, e.g., hair cream, hand cleaner, bath oil, suntan oil, brilliantine, anti-perspirants, perfumes and colognes, cold creams, electric pre-shave, eye and throat oil, fingernail polish, topical pharmaceutical ointments, lipsticks, stick rouge, skin lotions and creams, skin moisturizers, cleansing creams and after bath splash and lotions.

DETDESC:

DETD(220)

% by wt.

A. Beeswax, white	11.	00
Cetyl alcohol	2.50	
Cetyl palmitate	2.20	
Mineral oil, light, **	NF**	28.00
Isostearyl benzoate	20.6	0
Cerasynt Q (Glycery	l stearate,	self-emulsifying).sup.
	0.75	
Propyl paraben (prop	yl p-**h	(droxybenzoate**)
	0.05	
B. Water, purified	32.8	3
Borax (sodium borat	e) 0.1	75
Methyl paraben (met	hyl p-**h	ydroxybenzoate**)
	^`.;	

C. Water, purified 1.00 Dowicil 200 (Quaternium-15).sup.2 0.10

D. Fragrance

0.07

.sup.1 Van Dyk. . .

US PAT NO: 4,322,545 [IMAGE AVAILABLE] 1.4: 17 of 19 DATE ISSUED: Mar. 30, 1982 TITLE: Benzoic acid esters

Thomas L. Scala, Jr., West Milford, NJ INVENTOR: ASSIGNEE: Finetex, Inc., Elmwood Park, NJ (U.S. corp.)

APPL-NO: 06/252,794 DATE FILED: Apr. 13, 1981 ART-UNIT: 126 PRIM-EXMR: Paul J. Killos LEGAL-REP: Weingram & Klauber

US PAT NO: 4,322,545 [IMAGE AVAILABLE]

L4: 17 of 19

ABSTRACT:

A substantially pure benzoic acid ester of a mixture of alcohols. The mixture of alcohols consists essentially of (A) at least one C.sub.12 or C.sub.14 primary alcohol and (B) at least one C.sub.13 or C.sub.15 primary alcohol. The weight ratio of the even carbon number alcohols (A) to the odd carbon number alcohols (B) is from about 0.25:1 to about 4:1. preferably from about 0.5:1 to about 3:1. At least 70% by weight of each alcohol is linear and substantially all of the remainder of each alcohol is branched at the two carbon position. The compositions of this invention have unique properties in that they are substantially non-greasy, lack oiliness and greasiness, have low cloud points and pour points, have a bland odor, low toxicity, and are stable. The properties make such compositions useful as a vehicle or carrier, emollient or solubilizer for toiletry and cosmetic formulations, e.g., hair cream. hand cleaner, bath oil, suntan oil, brilliantine, anti-perspirants, perfumes and colognes, cold creams, electric pre-shave, eye and throat oil, fingernail polish, topical pharmaceutical ointments, lipsticks, stick rouge, skin lotions and creams, skin moisturizers, cleansing creams and after bath splash and lotions.

DETDESC:

DETD(334)

% by wt.

A. Beeswax, white	11.00
Cetyl alcohol	2.50
Cetyl palmitate	2.20
Mineral oil, light, **	NF** 28.00
NEODOL 25 benzoas	e 20.60
Cerasynt Q (Glyceryl	stearate, self-emulsifying).sup
0).75
Propyl paraben (prop	yl p-**hydroxybenzoate**)
	0.05
B. Water, purified	32.83
Borax (sodium borate) 0.75
Methyl paraben (meth	nyl p-**hydroxybenzoate**)
	0.15
C. Water, purified	1.00
Dowicil 200 (Quaterr	ium-15).sup.2
	0.10

.sup.1 Van Dyk. . .

D. Fragrance

```
US PAT NO: 4,306,076 [IMAGE AVAILABLE]
                                                L4: 18 of 19
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TITLE: Inter-phenylene CBA compounds INVENTOR: Norman A Marin Norman A. Nelson, Galesburg, MI

0.07

ASSIGNEE: The Upjohn Company, Kalamazoo, MI (U.S. corp.)

APPL-NO: 06/219,131 DATE FILED: Dec. 22, 1980 ART-UNIT: 126 PRIM-EXMR: Paul J. Killos

LEGAL-REP: L. Ruth Hattan, Robert A. Armitage

US PAT NO: 4,306,076 [IMAGE AVAILABLE] L4: 18 of 19

ABSTRACT:

The present specification provides novel analogs of carbacyclin (CBA.sub.2), 6a-carba-prostacyclin (6a-carba-PGI.sub.2), which have pronounced prostacyclin-like pharmacological activity, e.g., as platelet anti-aggregatory agents. Specifically the novel chemical analogs of CBA.sub.2 are those substituted by fluoro (C-5), alkyl (C-9), interphenylene (C-5), and methano (C-6a,9). Further provided are benzindene analogs of CBA.sub.2 and substituted forms thereof, i.e.. 9-deoxy-2',9-methano (or 2',9-methano)-3-oxa-4,5,6-trinor-3,7-(1',3'interphenylene)-PGF.sub.1 compounds. Also provided are a variety of novel chemical intermediates, e.g., substituted bicyclo[3.3.0]octane intermediates, and chemical process utilizing such intermediates which are useful in the preparation of the novel CBA.sub.2 analogs.

SUMMARY:

BSUM(132)

. N-methyl-N-cyclohexylamide, N-ethyl-N-cyclopentylamide, and N-ethyl-Ncyclohexylamide. Amides within the scope of aralkylamino are benzylamide, 2-phenylethylamide, and N-methyl-N benzyl-amide. Amides within the scope **nf** substituted phenylamide are p-chloroanilide m-chloroanilide, 2,4-dichloroanilide, 2,4,6-trichloroanilide, m-nitroanilide, p-nitroanilide, p-methoxyanilide, 3,4-dimethoxyanilide, 3,4,5-trimethoxyanilide, p-hydroxymethylanilide, p-methylanilide, m-methyl anilide, p-ethylanilide, t-butylanilide, p-carboxyanilide,

p-methoxycarbonyl. . . benzoylalkylamino are pchlorobenzoylmethylamide, m-chlorobenzoylmethylamide, 2,4-dichlorobenzoylmethylamide, 2,4,6-trichlorobenzoylmethylamide, m-nitrobenzoylmethylamide, p-nitrobenzoylmethylamide, p-methoxybenzoylmethylamide, 2,4-dimethoxy benzoylmethylamide, 3,4,5-trimethoxybenzoylmethylamide, p-hydroxymethylbenzoylmethylamide, p-methylbenzoylmethylamide, m-methylbenzoylmethylamide, p-ethylbenzoylmethylamide, t-butylbenzoylmethylamide, p-carboxybenzoylmethylamide, m-methoxycarbonylbenzoylmethylamide, o-carboxybenzoylmethylamide, o-**hydroxybenzoylmethylamide**, p-chlorobenzoylethylamide, m-chlorobenzoylethylamide, 2,4-dichlorobenzoylethylamide, 2,4,6-trichlorobenzoylethyla m-nitrobenzoylethylamide, p-nitrobenzoylethylamide, pmethoxybenzoylethylamide, p-methoxybenzoylethylamide, 2,4-dimethoxybenzoylethylamide, 3,4,5trimethoxybenzoylethylamide, p-hydroxymethylbenzoylethylamide, p-methylbenzoylethylamide, m-methylbenzoylethylamide, p-ethylbenzoylethylamide, tbutylbenzoylethylamide, p-carboxybenzoylethylamide, m-methoxycarbonylbenzoylethylamide, o-carboxybenzoylethylamide, o.**hydroxybenzoylethylamide**, p-chlorobenzoylpropylamide, m-chlorobenzoylpropylamide, 2,4-dichlorobenzoylpropylamide, 2,4,6-trichlorobenzoylpropylamide, m-nitrobenzoylpropylamide, p-nitrobenzoylpropylamide, p-methoxybenzoylpropylamide, 2,4-dimethoxybenzoylpropylamide, 3,4,5-trimethoxybenzoylpropylamide, p-hydroxymethylbenzoylpropylamide, p-methylbenzoylpropylamide, m-methylbenzoylpropylamide, p-ethylbenzoylpropylamide, t-butylbenzoylpropylamide, p-carboxybenzoylpropylamide, m-methoxycarbonylbenzoylpropylamide, o-carboxybenzoylpropylamide, o-arboxybenzoylpropylamide, m-chlorobenzoylbutylamide, 2,4-dichlorobenzoylbutylamide, 2,4,6-trichlorobenzoylbutylamide, m-nitrobenzoylmethylamide, p-nitrobenzoylbutylamide, p-methoxybenzoylbutylamine, 2,4-dimethoxybenzoylbutyl-amide, 3,4,5-trimethoxybenzoylbutylamide, p-hydroxymethylbenzoylbutyl-amide, p-methylbenzoylbutyamide, m-methylbenzoylbutylamide, p-ethyl-benzoylbutylamide, m-methylbenzoylbutylamide, p-ethylbenzoylbutyl-amide, t-butylbenzoylbutylamide, p-carboxybenzoylbutylamide, m-methoxycarbonylbenzoylbutylamide, o-carboxybenzoylbutylamide, o-**hydroxybenzoylmethylamide**. Amides within the scope of pyridylamino are alpha-pyridylamide, .beta-pyridylamide, and .gamma.-pyridylamide. Amides within the scope of substituted pyridylamino are 4-methyl-alpha.-pyridylamide, . .

US PAT NO: 4,306,075 [IMAGE AVAILABLE] DATE ISSUED: Dec. 15, 1981

L4: 19 of 19

L4: 19 of 19

TITLE: Composition and process

INVENTOR: Paul A. Aristoff, Portage, MI

The Upjohn Company, Kalamazoo, MI (U.S. corp.) ASSIGNEE:

APPL-NO: 06/219,210 DATE FILED: Dec. 22, 1980 ART-UNIT: 126

PRIM-EXMR: PauL J. Killos

LEGAL-REP: L. Ruth Hattan, Robert A. Armitage

US PAT NO: 4,306,075 [IMAGE AVAILABLE]

The present specification provides novel analogs of carbacyclin (CBA.sub.2), 6a-carba-prostacyclin (6a-carba-PGI.sub.2), which have pronounced prostacyclin-like pharmacological activity, e.g., as platelet antiaggregatory agents. Specifically the novel chemical analogs of CBA.sub.2 are those substituted by fluoro (C-5), alkyl (C-9), interphenylene (C-5), and methano (C-6a,9). Further provided are benzindene analogs of CBA.sub.2 and substituted forms thereof, i.e., 9-deoxy-2',9-methano (or 2',9-metheno)-3-oxa-4,5,6-trinor-3,7-(1',3'-interphenylene)-PGF.sub.1 compounds. Also provided are a variety of novel chemical intermediates, e.g., substituted bicyclo[3.3.0]octane intermediates, and chemical process utilizing such intermediates which are useful in the preparation of the novel CBA.sub.2 analogs.

SUMMARY:

BSUM(132)

. N-methyl-N-cyclohexylamide, N-ethyl-N-cyclopentylamide, and N-ethyl-Ncyclohexylamide. Amides within the scope of aralkylamino are benzylamide, 2-phenylethylamide, and N-methyl-N benzyl-amide. Amides within the scope "*nf" substituted phenylamide are p-chloroanilide, m-chloroanilide, 2,4-dichloroanilide, 2,4-6-trichloroanilide, m-nitroanilide, p-nitroanilide, p-methoxyanilide, 3,4-dimethoxyanilide, 3,4,5-trimethoxyanilide, p-hydroxymethylanilide, p-methylanilide, m-methyl anilide, p-ethylanilide, t-butylanilide, p-carboxyanilide, p-methoxycarbonyl. . . benzoylalkylamino are pchlorobenzoylmethylamide, m-chlorobenzoylmethylamide, 2,4-dichlorobenzoylmethylamide, 2,4,6-trichlorobenzoylmethylamide, m-nitrobenzoylmethylamide, p-nitrobenzoylmethylamide, p-methoxybenzoylmethylamide, 2,4-dimethoxy benzoylmethylamide, 3,4,5-trimethoxybenzoylmethylamide, p-hydroxymethylbenzoylmethylamide, p-methylbenzoylmethylamide, m-methylbenzoylmethylamide,

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p-ethylbenzoylmethylamide, t-butylbenzoylmethylamide,
p-carboxybenzoylmethylamide, m-methoxycarbonylbenzoylmethylamide, o-carboxybenzoylmethylamide, o-**hydroxybenzoylmethylamide**,
p-chlorobenzoylethylamide, m-chlorobenzoylethylamide,
 2,4-dichlorobenzoylethylamide, 2,4,6-trichlorobenzoylethylamide,
m-nitrobenzoylethylamide, p-nitrobenzoylethylamide, p-methoxybenzoylethylamide, p-methoxybenzoylethylamide, p-methoxybenzoylethylamide, 2,4-dimethoxybenzoylethylamide, 3,4,5trimethoxybenzoylethylamide,
p-hydroxymethylbenzoylethylamide, p-methylbenzoylethylamide,
m-methylbenzoylethylamide, p-ethylbenzoylethylamide, t-
butylbenzoylethylamide, p-carboxybenzoylethylamide, m-
omethoxycarbonybenzoylethylamide, o-carboxybenzoylethylamide,
o-*hydroxybenzoylethylamide**, p-chlorobenzoylpropylamide,
m-chlorobenzoylpropylamide, 2,4-dichlorobenzoylpropylamide,
2,4,6-trichlorobenzoylpropylamide, m-nitrobenzoylpropylamide
p-nitrobenzoylpropylamide, p-methoxybenzoylpropylamide,
2,4-dimethoxybenzoylpropylamide, 3,4,5-trimethoxybenzoylpropylamide,
p-hydroxymethylbenzoylpropylamide, p-methylbenzoylpropylamide,
 m-methylbenzoylpropylamide, p-ethylbenzoylpropylamide,
t-butylbenzoylpropylamide, p-carboxybenzoylpropylamide, m-methoxycarbonylbenzoylpropylamide, o-carboxybenzoylpropylamide, o-*hydroxybenzoylpropylamide**, p-chlorobenzoylbutylamide,
o--nydroxybenzoylpropylamide--, p-cniorobenzoylbutylamide,
m-chlorobenzoylbutylamide, 2,4-dichlorobenzoylbutylamide,
2,4.6-trichlorobenzoylbutylamide, m-nitrobenzoylmethylamide,
p-nitrobenzoylbutylamide, p-methoxybenzoylbutylamine,
2,4-dimethoxybenzoylbutyl-amide, 3,4,5-trimethoxybenzoylbutylamide,
p-hydroxymethylbenzoylbutyl-amide, p-methylbenzoylbutyamide,
m-methylbenzoylbutylamide, p-ethyl-benzoylbutylamide,
m-methylbenzoylbutylamide, p-ethylbenzoylbutyl-amide,
t-butylbenzoylbutylamide, p-carboxybenzoylbutylamide,
m-methoxycarbonylbenzoylbutylamide, o-carboxybenzoylbutylamide, o-**hydroxybenzoylmethylamide**. Amides within the scope of pyridylamino
are .alpha.-pyridylamide, .beta.-pyridylamide, and .gamma.-pyridylamide.
Amides within the scope of substituted pyridylamino are 4-methyl-.alpha.-pyridylamide,. . .
 => s ((nuclear factor) (3a) (b or K or kappa)) or (nfkb) or (nf kb)
           62536 NUCLEAR
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268223 FACTOR
 205 NUCLEAR FACTOR
     (NUCLEAR(W)FACTOR)
1219215 B
374077 K
 4991 KAPPA
  59 (NUCLEAR FACTOR) (3A) (B OR K OR KAPPA)
51 NFKB
 11097 NF
 16285 KB
 118 NF KB
     (NF(W)KB)
   198 ((NUCLEAR FACTOR) (3A) (B OR K OR KAPPA)) OR (NFKB) OR
```

(NF KB)

= > s 12 (p) 15

1.6 0 L2 (P) L5

= > s 12 and 15

10 L2 AND L5

0 LI AND L7 1.8

= > d 17 1-10 bib rel ab

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US PAT NO: 5,885,829 [IMAGE AVAILABLE]
                                                                 L7: 1 of 10
DATE ISSUED: Mar. 23, 1999
TITLE: Engineering oral tissues
INVENTOR: David J. Mooney, Ann Arbor, MI
Robert B. Rutherford, Ann Arbor, MI
ASSIGNEE:
                 The Regents of the University of Michigan, Ann Arbor, MI
(U.S. corp.)
APPL-NO: 08/864,494
DATE FILED: May 28, 1997
ART-UNIT: 191
PRIM-EXMR: Na
                  Nancy Degen
Arnold, White & Durkee
LEGAL-REP:
US PAT NO: 5,885,829 [IMAGE AVAILABLE]
                                                                L7: 1 of 10
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Disclosed are methods for regenerating dental and oral tissues from viable cells using ex vivo culture on a structural matrix. The regenerated oral tissues and tissue-matrix preparations thus provided have both clinical applications in dentistry and oral medicine and are

also useful in in vitro toxicity and biocompatibility testing. US PAT NO: 5,874,448 [IMAGE AVAILABLE] DATE ISSUED: Feb. 23, 1999 L7: 2 of 10 Substituted 2-(2,6 dioxo-3-fluoropiperidin-3-yl)-isoindolines and method of reducing TNF alpha. levels George W. Muller, Bridgewater, NJ TITLE: INVENTOR: David I. Stirling, Branchburg, NJ Roger Shen-Chu Chen, Edison, NJ Hon-Wah Man, Neshanic Station, NJ ASSIGNEE: Celgene Corporation, Warren, NJ (U.S. corp.) 08/976,140 APPL-NO: DATE FILED: Nov. 18, 1997 ART-UNIT: 162 PRIM-EXMR: Evelyn Huang Mathews, Collins, Shepherd & Gould, P.A. LEGAL-REP: US PAT NO: 5,874,448 [IMAGE AVAILABLE] L7: 2 of 10 ABSTRACT: 1-Oxo-2-(2,6-dioxo-3-fluoropiperidin-3-yl)isoindolines and 1,3-dioxo-2-(2,6-dioxo-3-fluoropiperidin-3-yl)isoindolines reduce the levels of TNF.alpha. in a mammal. A typical embodiment is 1,3-dioxo-2-(2,6-dioxo-3-fluoropiperidin-3-yl)-isoindoline. US PAT NO: 5,849,263 [IMAGE AVAILABLE] DATE ISSUED: Dec. 15, 1998 L7: 3 of 10 Pharmaceutical compositions containing alkylaryl polyether alcohol polymer
R: Thomas P. Kennedy, Richmond, VA INVENTOR: Charlotte-Mecklenburg Hospital Authority, Charlotte, NC ASSIGNEE: (U.S. corp.) APPL-NO: 08/638,893 DATE FILED: Apr. 25, 1996 166 ART-UNIT: PRIM-EXMR: Robert H. Harrison Bell Seltzer Intellectual Property Law Group of Alston & LEGAL-REP: Bird LLP US PAT NO: 5,849,263 [IMAGE AVAILABLE] L7: 3 of 10 REL-US-DATA: Continuation-in-part of Ser. No. 299,316, Aug. 31, 1994, Pat. No. 5,512,270, which is a continuation-in-part of Ser. No. 39,732, Mar. 30, 1993, abandoned. There is provided novel pharmaceutical compositions containing tyloxapol

There is provided novel pharmaceutical compositions containing tyloxapol as the active ingredient. These formulations comprise tyloxapol at concentrations above 0.125%, preferably from about 0.25% to about 5.0%. In addition, the invention encompasses pharmaceutical compositions having reduced hypertonicity which compositions comprise tyloxapol in pharmaceutically acceptable solutions without significant concentrations of hypertonic agents or other active ingredients NaHCO.sub.3, or active phospholipids, such as DPPC. The less hypertonic formulations allow one to derive all the benefits of the active ingredient tyloxapol, such as its reduced toxicity and enhanced half-life, while avoiding or reducing side effects, such as bronchospasms, associated with the various hypertonic agents or other active ingredient agents.

US PAT NO: 5,840,277 [IMAGE AVAILABLE] L7: 4 of 10
DATE ISSUED: Nov. 24, 1998
TITLE: Treatment of chronic pulmonary inflammation
INVENTOR: Andrew J. Ghio, Durham, NC
Thomas P. Kennedy, Richmond, VA
ASSIGNEE: Apr. Charlotte Hospital Authority, Charlotte, NC (U.S. corp.)
08/632,275
DATE FILED: Apr. 15, 1996
ART-UNIT: 185
PRIM-EXMR: David Guzo
LEGAL-REP: The Bell Seltzer Intellectual Law Firm of Alston & Bird, LLP

US PAT NO: 5,840,277 [IMAGE AVAILABLE] L7: 4 of 10 REL-US-DATA: Continuation-in-part of Ser. No. 413,699, Mar. 30, 1995, which is a continuation-in-part of Ser. No. 219,770, Mar. 29, 1994, Pat. No. 5,474,760, Dec. 12, 1995, which is a continuation-in-part of Ser. No. 299,316, Aug. 31, 1994, Pat. No. 5,512,270, which is a continuation-in-part of Ser. No. 39,732, Mar. 30, 1993, abandoned

ABSTRACT

A method and medicant for the inhibition of activation of the nuclear transcription NF- kappa. B comprising administering an effective amount of a compound of the formula: ##STR!## where R=ethylene, R'=C.sub.4 to C.sub.14 straight chain or branched alkyl, x is greater than 1, and y=8 to 18 is provided. The medicant is preferably administered by aerosolization into the mammalian respiratory system. The medicant may

also be applied to the mammalian skin. Preferably the medicant includes a physiologically acceptable carrier which may be selected from buffered saline, isotonic saline, normal saline, petroleum-based ointments and U.S.P. cold cream. There is further provided a method wherein said medicant includes an anti-inflammatory steroid. In addition a method and medicant for treating cutaneous inflammatory disorders, inhibiting the secretion of the pro-inflammatory cytokines TNF, IL-1, IL-6, IL-8 and the growth factor GM-CSF is provided.

US PAT NO: 5,798,368 [IMAGE AVAILABLE]
DATE ISSUED: Aug. 25, 1998 L7: 5 of 10 Tetrasubstituted 2-(2,6-dioxopiperidin-3-yl)-1oxoisoindolines and method of reducing TNF.alpha. levels DR: George W. Muller, Bridgewater, NJ David I. Stirling, Branchburg, NJ Roger Shen-Chu Chen, Edison, NJ INVENTOR: ASSIGNEE: Celgene Corporation, Warren, NJ (U.S. corp.) 08/701 494 APPL-NO: DATE FILED: Aug. 22, 1996 ART-UNIT: 164 PRIM-EXMR: James H. Reamer Mathews, Collins, Shepherd & Gould, P.A. LEGAL-REP: US PAT NO: 5,798,368 [IMAGE AVAILABLE] L7: 5 of 10

ABSTRACT:

Tetrasubstituted 1-oxo-2-(2,6-dioxopiperidin-3-yl)isoindolines reduce the levels of TNF.alpha. in a mammal. A typical embodiment is 1-oxo-2-(2,6-dioxopiperidin-3-yl)-4,5,6,7-tetrafluoroisoindoline.

US PAT NO: 5,670,617 [IMAGE AVAILABLE]
DATE ISSUED: Sep. 23, 1997
TITLE: Nucleic acid conjugates of tat-derived transport L7: 6 of 10 polypeptides Carl Pabo, 18 Weldon Rd., Newton, MA 02158

James G. Barsoum, 9 Marlboro Rd., Lexington, MA 02173 INVENTOR: Stephen E. Fawell, One Black Horse Ter., Winchester, MA 01890 R. Blake Pepinsky, 30 Falmouth Rd., Arlington, MA 02174 08/450,246 APPL-NO: DATE FILED: May 25, 1995 ART-UNIT: 189 PRIM-EXMR: George C. Elliot ASST-EXMR: Thomas G. Larson LEGAL-REP: James F. Haley, Jr., Madge R. Kanter

US PAT NO: 5,670,617 [IMAGE AVAILABLE] L7: 6 of 10 REL-US-DATA: Division of Ser. No. 235,403, Apr. 28, 1994, which is a continuation-in-part of Ser. No. 158,015, Nov. 24, 1993, abandoned, which is a continuation of Ser. No. 636,662, Jan. 2, 1991, abandoned, which is a continuation-in-part of Ser. No. 454,450, Dec. 21, 1989, abandoned, said Ser. No. 235,403 is a continuation-in-part of Ser. No. 934,375, Aug. 21, 1992, abandoned.

ABSTRACT:

This invention relates to delivery of biologically active cargo molecules, such as polypeptides and nucleic acids, into the cytoplasm and nuclei of cells in vitro and in vivo. Intracellular delivery of cargo molecules according to this invention is accomplished by the use of novel transport polypeptides which comprise HIV tat protein or one or more portions thereof, and which are covalently attached to cargo molecules. The transport polypeptides in preferred embodiments of this invention are characterized by the presence of the tat basic region (amino acids 49-57), the absence of the tat cysteine-rich region (amino acids 22-36) and the absence of the tat exon 2-encoded carboxy-terminal domain (amino acids 73-86) of the naturally-occurring tat protein. By virtue of the absence of the cysteine-rich region, the preferred transport polypeptides of this invention solve the potential problems of spurious trans-activation and disulfide aggregation. The reduced size of the preferred transport polypeptides of this invention also minimizes interference with the biological activity of the cargo molecule.

US PAT NO: 5,635,517 [IMAGE AVAILABLE] L7: 7 of 10
DATE ISSUED: Jun. 3, 1997
TITLE: Method of reducing TNF.alpha. levels with amino substituted 2-(2,6-dioxopiperidin-3-yl)-1-oxo-and 1,3-dioxosiondolines
INVENTOR: George W. Muller, Bridgewater, NJ David I. Stirling, Branchburg, NJ Roger S. -C. Chen, Edison, NJ
ASSIGNEE: Celgene Corporation, Warren, NJ (U.S. corp.)
APPL-NO: 08/690,258
DATE FILED: Jul. 24, 1996
ART-UNIT: 123
PRIM-EXMR: C. Warren Ivy

ASST-EXMR: C. S. Aulakh

Mathews, Collins, Shepherd & Gould, P.A. LEGAL-REP:

US PAT NO: 5,635,517 [IMAGE AVAILABLE]

L7: 7 of 10

ABSTRACT:

1-Oxo- and 1,3-dioxo-2-(2,6-dioxopiperidin-3-yl) isoindolines substituted with amino in the benzo ring reduce the levels of TNF alpha. in a mammal. A typical embodiment is 1,3-dioxo-2-(2,6-dioxopiperidin-3-yl)-5aminoisoindoline.

US PAT NO: 5,612,330 [IMAGE AVAILABLE] DATE ISSUED: Mar. 18, 1997

L7: 8 of 10

TITLE: Methods for inhibiting and controlling viral growth INVENTOR: David T. Connor, Ann Arbor, MI

Stephen J. Gracheck, Ann Arbor, MI

Warner-Lambert Company, Morris Plains, NJ (U.S. corp.) ASSIGNEE:

08/408,431 APPL-NO: DATE FILED: Mar. 22, 1995

122

PRIM-EXMR: Robert T. Bond

LEGAL-REP: Charles W. Ashbrook

US PAT NO: 5,612,330 [IMAGE AVAILABLE] L7: 8 of 10 REL-US-DATA: Continuation-in-part of Ser. No. 351,611, Dec. 12, 1994, Pat. No. 5,489,586, which is a continuation-in-part of Ser. No. 207,330, Mar. 7, 1994, abandoned.

ABSTRACT:

Benzothiophene, benzofuran and indolethiazepinones, oxazepinones, and diazepinones are effective therapeutic agents for treating viral diseases, including those caused by herpesvirus and HIV.

US PAT NO: 5,608,095 [IMAGE AVAILABLE]

L7: 9 of 10

DATE ISSUED: Mar. 4, 1997

Alkyl-4-silyl-phenols and esters thereof as antiatherosclerotic agents TITLE:

INVENTOR: Roger A. Parker, Cincinnati, OH

Michael L. Edwards, Cincinnati, OH Mark J. Vaal, Cincinnati, OH James E. Matt, Jr., Cincinnati, OH

Kim S. Chen, Cincinnati, OH Mark T. Yates, Ann Arbor, Ml Paul S. Wright, Cincinnati, OH

E: Hoechst Marion Roussel, Inc., Cincinnati, OH (U.S. corp.) ASSIGNEE:

08/637.968 APPL-NO:

DATE FILED: Apr. 30, 1996

ART-UNIT: 124

PRIM-EXMR: Paul F. Shaver

LEGAL-REP: William R. Boudreaux

US PAT NO: 5,608,095 [IMAGE AVAILABLE] L7: 9 of 10

ABSTRACT:

This invention relates to compounds of the formula ##STR1## wherein R.sub.1 and R.sub.6 are each independently C.sub.1 -C.sub.6 alkyl; R.sub.2, R.sub.3 and R.sub.4 are each independently hydrogen or C.sub.1 -C.sub.6 alkyl;

R is hydrogen or --C(O)--(CH.sub.2).sub.m --Q wherein Q is hydrogen or --COOH and m is an integer 1, 2, 3 or 4;

Z is a thio, oxy or methylene group;
A is a C.sub.1 -C.sub.4 alkylene group;
R.sub.5 and R.sub.7 are each independently a C.sub.1 -C.sub.6 alkyl or --(CH.sub.2).sub.n --(Ar) wherein n is an integer 0, 1, 2 or 3; and Ar is phenyl or naphthyl unsubstituted or substituted with one to three is pitely to liaghtly instantiated of substituted with the substitutents selected from the group consisting of hydroxy, methoxy, ethoxy, halogen, trifluoromethyl, C.sub.1 -C.sub.6 alkyl, or --NR.sub.8 R.sub.9, wherein R.sub.8 and R.sub.9 are each independently hydrogen or C.sub.1 -C.sub.6 alkyl; with the proviso that when R.sub.2 and at least one of R.sub.5 or R.sub.7 is C.sub.1 -C.sub.6 alkyl, and Ar is not substituted with trifluoromethyl or --NR.sub.8 R.sub.9, then R is --C(O)--(CH.sub.2).sub.m --Q; or a pharmaceutically acceptable salt thereof; useful for the treatment of atherosclerosis and chronic inflammatory disorders; for inhibiting cytokine-induced expression of VCAM-1 and/or ICAM-1; for inhibiting the peroxidation of LDL lipid; for lowering plasma cholesterol; and as antioxidant chemical additives useful for preventing oxidative deterioration in organic materials.

US PAT NO: 5,317,019 [IMAGE AVAILABLE]

L7: 10 of 10

DATE ISSUED: May 31, 1994

Inhibition of interleukin-1 and tumor necrosis factor

production by monocytes and/or macrophages

INVENTOR: Paul E. Bender, Cherry Hill, NJ

Don E. Griswold, North Wales, PA Nabil Hanna, Solana Beach, CA

John C. Lee, Radnor, PA

Bartholomew J. Votta, Pottstown, PA Philip L. Simon, Randolph, NJ Alison M. Badger, Bryn Mawr, PA

Klaus M. Esser, Downingtown, PA

SmithKline Beecham Corp., Phildelphia, PA (U.S. corp.) ASSIGNEE:

07/809,484 APPL-NO: DATE FILED: Dec. 12, 1991

ART-UNIT: 123

C. Warren Ivy PRIM-EXMR:

ASST-EXMR: Raymond Covington

LEGAL-REP: Dara L. Dinner, Stephen Venetianer, Edward T. Lentz

US PAT NO: 5,317,019 [IMAGE AVAILABLE] L7: 10 of 10 REL-US-DATA: Continuation-in-part of Ser. No. 365,349, Jun. 13, 1989, abandoned.

ABSTRACT:

A method of inhibiting the production of interleukin-1 by monocytes and/or macrophages in a human in need thereof which comprises administering to such a human an effective, interleukin-1 production inhibiting amount of a diaryl-substituted imidazole fused to a second heterocyclic ring containing a nitrogen bridgehead atom wherein said second ring may also contain sulfur, oxygen or an additional nitrogen atom, and may contain additional unsaturation.

This invention relates to a method of inhibiting the production of Tumor Necrosis Factor (TNF) by monocytes or macrophages in a human in need thereof which comprises administering to such mammal an effective, TNF production inhibiting amount of a compound of Formula (I) as described herein. The compounds of Formula (II) are generally described as diaryl-substituted imidazole fused to a second heterocyclic ring containing a nitrogen bridgehead wherein said ring may also contain sulfur, oxygen, or an additional nitrogen atom, and may contain additional unsaturation.

=> s (12 or 15) (p) (treat? or therap? or medic? or pharmac? or drug#)

586240 TREAT? 89576 THERAP?

142287 MEDIC? 116190 PHARMAC?

73018 DRUG#

6200 (L2 OR L5) (P) (TREAT? OR THERAP? OR MEDIC? OR PHARMAC? OR DRUG#

= > s 19 and 514/clas

80898 514/CLAS

3942 L9 AND 514/CLAS L10

= > s 19 and 564/clas

32158 564/CLAS

471 L9 AND 564/CLAS

=> s 110 and 111

288 L10 AND L11 L12

= > s 15 and 514/clas

80898 514/CLAS 72 L5 AND 514/CLAS L13

= > s 15 and 564/clas

32158 564/CLAS

6 L5 AND 564/CLAS

=> s 113 or 114

72 L13 OR L14

= > d 1-72 bib kwic

US PAT NO: 5,905,089 [IMAGE AVAILABLE]

L15: 1 of 72

DATE ISSUED: May 18, 1999

Use of sesquiterpene lactones for treatment of severe

inflammatory disorders
INVENTOR: Daniel H. Hwang, Baton Rouge, LA
Nikolaus H. Fischer, Baton Rouge, LA

ASSIGNEE: Board of Supervisors of Louisiana State University and Agricultural and Mechanical College, Baton Rouge, LA

(U.S. corp.) 09/059,480 APPL-NO:

DATE FILED: Apr. 13, 1998

ART-UNIT: 164
PRIM-EXMR: Keith D. MacMillan

LEGAL-REP: Bonnie J. Davis, John H. Runnels

US PAT NO: 5,905,089 [IMAGE AVAILABLE] US-CL-CURRENT: **514/468**

L15: 1 of 72

L15: 2 of 72

DETDESC

DETD(60)

Parthenolide inhibits Nuclear Factor--kB (**NF**-**kB**) transcription factor that has been activated by LPS in the murine macrophage cell line (RAW 264.7) as assessed by the degradation of the protein, IKB.sub..alpha.. Activated **NF**-**kB** is known to induce the expression of many early response genes including inducible nitric oxide synthetase, cyclooxygenase, and chemokines that. . .

US PAT NO: 5,900,430 [IMAGE AVAILABLE] DATE ISSUED: May 4, 1999

ITILE: Cytokine inhibitors
INVENTOR: Alison Mary Badger, Bryn Mawr, PA
Wanda Bernadette High, Wayne, PA
ASSIGNEE: AnorMED, Inc., Langley, Canada (foreign corp.)

APPL-NO: 08/779,418 DATE FILED: Jan. 7, 1997 ART-UNIT: 125 PRIM-EXMR: Jerome D. G

PRIM-EXMR: Jerome D. Goldberg
LEGAL-REP: Kate H.Morrison & Foerster, LLP Murashige

US PAT NO: 5,900,430 [IMAGE AVAILABLE] 1.15: 2 of 72

US-CL-CURRENT: **514/409**, **212**, **278**

DETDESC:

DETD(29)

Osborn . . . molecular mechanism for the virus inducing activity of TNF is due to TNF's ability to activate a gene regulatory protein (**NF**-**kB**) found in the cytoplasm of cells, which promotes HIV replication through binding to a viral regulatory gene sequence (LTR).

US PAT NO: 5,883,081 [IMAGE AVAILABLE]
DATE ISSUED: Mar. 16, 1999

L15: 3 of 72

L15: 4 of 72

Isolation of novel HIV-2 proviruses TITLE

INVENTOR: Gunter Kraus, La Jolla, CA Flossie Wong-Staal, San Diego, CA Randy Talbott, Princeton, NJ

Eric M. Poeschla, San Diego, CA

The Regents of the University of California, Oakland, CA ASSIGNEE:

(U.S. corp.) 08/659,251 APPL-NO:

DATE FILED: Jun. 7, 1996

168 ART-UNIT:

PRIM-EXMR: Jeffrey Stucker

ASST-EXMR: Hankvel Park

LEGAL-REP: Townsend and Townsend and Crew

US PAT NO: 5,883,081 [IMAGE AVAILABLE] US-CL-CURRENT: **514/44**; 424/160.1; 435/69.1, 320.1; 530/388.35; 536/23.1

DETDESC:

DETD(215)

before the Spl binding sites. This deletion is not seen in other HIV-2 isolates, and is not similar to the **NFkB** duplication (Novembre, et al. (1991) Journal of Medical Primatology 20, 188-92) previously described in the SIV.sub.MMpbj LTR.

US PAT NO: 5,877,203 [IMAGE AVAILABLE]

DATE ISSUED: Mar. 2, 1999

Treatment for atherosclerosis and other cardiovascular and TITLE:

inflammatory diseases
INVENTOR: Russell M. Medford, Atlanta, GA

Margaret K. Offermann, Atlanta, GA

R. Wayne Alexander, Atlanta, GA
E: Emory University, Atlanta, GA (U.S. corp.)
c: 08/722,438 ASSIGNEE:

APPL-NO:

DATE FILED: Oct. 17, 1996

ART-UNIT: 163 PRIM-EXMR: Del

Deborah C. Lambkin

Sherry M. Knowles, JacquelineKing & Spalding Haley LEGAL-REP:

5,877,203 [IMAGE AVAILABLE] US PAT NO: US-CL-CURRENT: **514/423**, **212**, **330**, **551**, **599**, **707**, **712**

DETDESC:

DETD(28)

At the molecular level, PDTC has been shown to inhibit the activation of the transcriptional regulatory factor **Nf**-**kB** in response to certain cytokine and non-cytokine stimuli (Schreck, Rieber et al. 1991; Schreck, Meier et al. 1992). However, by. . . has been discovered that endothelial cells activate VCAM-I gene expression through an apparently novel transcriptional regulatory factor that is not "N(**.**kB**. This suggests that PDTC may regulate endothelial cell gene expression through its effect on a new transcriptional regulatory protein. It.

US PAT NO: 5,877,200 [IMAGE AVAILABLE] DATE ISSUED: Mar. 2, 1999

L15: 5 of 72

TITLE: Cyclic amides INVENTOR: George W.

yelic amides
George W. Muller, Bridgewater, NJ
Celgene Corporation, Warren, NJ (U.S. corp.) ASSIGNEE:

APPL-NO: 08/920,715 DATE FILED: Aug. 29, 1997 ART-UNIT: 162

PRIM-EXMR: John Kight

D. Margaret M. Mach ASST-EXMR:

Mathews, Collins Shepherd & Gould, P.A. LEGAL-REP:

US PAT NO: 5,877,200 [IMAGE AVAILABLE] US-CL-CURRENT: **514/411**; 548/450, 451

L15: 5 of 72

L15: 7 of 72

SUMMARY:

BSUM(13)

The **nuclear** **factor** .**kappa**.**B** (NF.**kappa**.B) is a pleiotropic transcriptional activator (Lenardo, et al. Cell 1989, 58, 227-29). NF kappa. B has been implicated as a transcriptional activator.

US PAT NO: 5,874,448 [IMAGE AVAILABLE]
DATE ISSUED: Feb. 23, 1999

Substituted 2-(2,6 dioxo-3-fluoropiperidin-3-yl)-

isoindolines and method of reducing TNF, alpha. levels R: George W. Muller, Bridgewater, NJ INVENTOR:

David I. Stirling, Branchburg, NJ

Roger Shen-Chu Chen, Edison, NJ

Hon-Wah Man, Neshanic Station, NJ

Celgene Corporation, Warren, NJ (U.S. corp.) ASSIGNEE:

08/976,140 APPL-NO: DATE FILED: Nov. 18, 1997

ART-UNIT: 162 PRIM-EXMR: Evelyn Huang

LEGAL-REP: Mathews, Collins, Shepherd & Gould, P.A.

US PAT NO: 5,874,448 [IMAGE AVAILABLE] US-CL-CURRENT: **514/323**; 546/201 L15: 6 of 72

BSUM(13)

The **nuclear** **factor** .**kappa**.**B** (NF.**kappa**.B) is a pleiotropic transcriptional activator (Lenardo, et al., Cell 1989, 58, 227-29). NF.kappa.B has been implicated as a transcriptional activator.

US PAT NO: 5,869,055 [IMAGE AVAILABLE]
DATE ISSUED: Feb. 9, 1999

ITITLE: Anti-inflammatory CD14 polypeptides INVENTOR: Shao-Chieh Juan, Moorpark, CA Henri S. Lichenstein, Boulder, CO

Samuel D. Wright, Westfield, NJ

Amgen, Inc., Thousand Oaks, CA (U.S. corp.) ASSIGNEE:

APPL-NO: 08/484,397 DATE FILED: Jun. 7, 1995

ART-UNIT: 186 PRIM-EXMR: Th

Thomas M. Cunningham

ASST-EXMR:

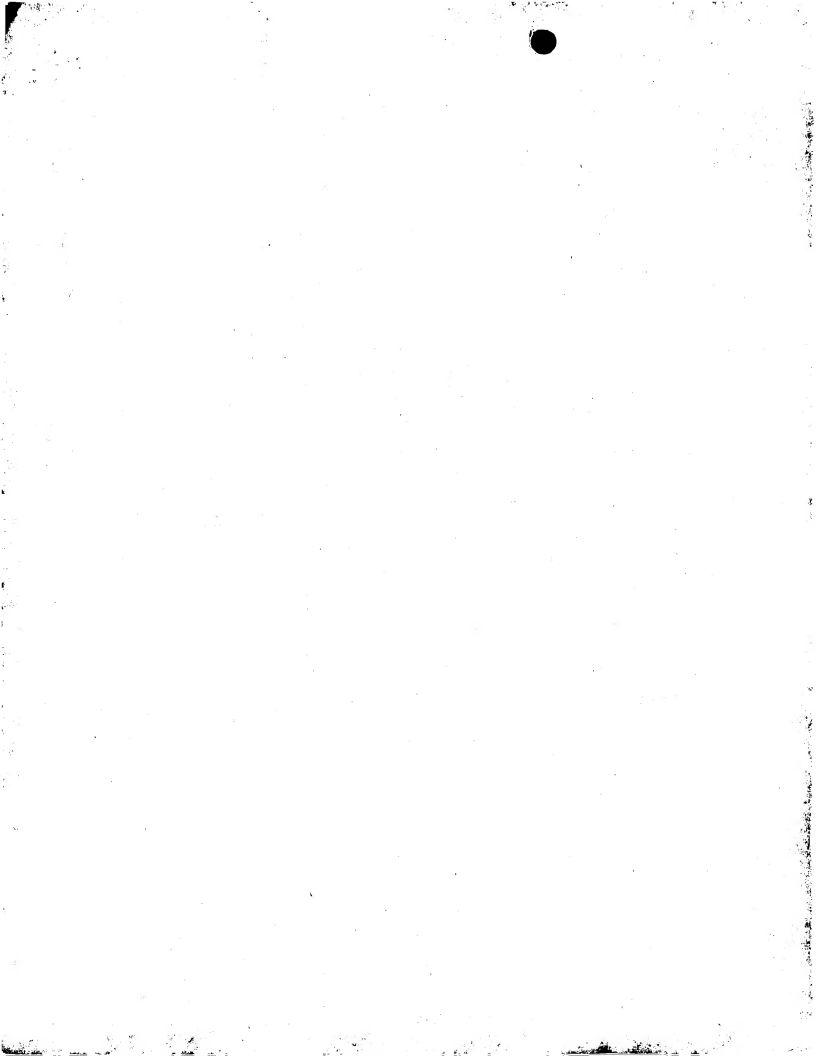
Martha T. Lubet Timothy J. Gaul, Ron K. Levy, Steven M. Odre LEGAL-REP:

US PAT NO: 5,869,055 [IMAGE AVAILABLE] L15: 7 of 72 US-CL-CURRENT: 424/185.1; **514/2**; 530/300, 317, 351; 536/23.5

DETDESC:

DETD(93)

LPS . . . eliminated formation of both complexes (data not shown). Stimulation of U373 cells with sCD14.sub.(7-10)A and LPS caused only 5%



of **NF**-**kB** activation as quantitated by gel scanning (FIG. 6, lane 6). Comparatively, stimulation of U373 cells with a mutant which does. .

US PAT NO: 5,863,904 [IMAGE AVAILABLE] DATE ISSUED: Jan. 26, 1999

TITLE: Methods for treating cancers and restenosis with P21
INVENTOR: Gary J. Nabel, Ann Arbor, MI
Zhi-yong Yang, Ann Arbor, MI

Elizabeth G. Nabel, Ann Arbor, MI

E: The University of Michigan, Ann Arbor, MI (U.S. corp.) ASSIGNEE:

08/533,942 APPL-NO: DATE FILED: Sep. 26, 1995

ART-UNIT: 189

PRIM-EXMR: Jasemine C. Chambers Karen M. Hauda ASST-EXMR: Brinks Hofer Gilson & Lione LEGAL-REP:

US PAT NO: 5,863,904 [IMAGE AVAILABLE] US-CL-CURRENT: **514/44**; 435/69.1, 375 L15: 8 of 72

SUMMARY:

BSUM(13)

. . as well as the induction of the differentiated phenotype arises from altered patterns of gene expression, mediated in part by **NF**-**kB**, resulting from p21 induced transcriptional regulation leading to terminal differentiation and growth arrest. Previous attempts to induce antitumor effects through. . .

US PAT NO: 5,861,290 [IMAGE AVAILABLE] DATE ISSUED: Jan. 19, 1999

L15: 9 of 72

L15: 8 of 72

Methods and polynucleotide constructs for treating host

cells for infection or hyperproliferative disorders
INVENTOR: Mark A. Goldsmith, 20 Maple St., West Roxbury, MA 02132

Robert O. Ralston, 2863 Judah, San Francisco, CA 94122

APPL-NO: 07/965,039

APYL-PNO: 07/903,039

DATE FILED: Oct. 22, 1992
ART-UNIT: 185

PRIM-EXMR: Jonny F. Railey, II

LEGAL-REP: Norman J. Kruse, Donald J. Pochopien, Robert P. Blackburn

US PAT NO: 5,861,290 [IMAGE AVAILABLE] L15: 9 of 72 US-CL-CURRENT: 424/93.2; 435/320.1; **514/44**; 536/23.1, 23.2, 23.5, 23.53, 23.6, 23.7, 23.72, 24.1, 24.5

DETDESC:

DETD(6)

The . . . contains both the tar region, which is highly selective for HIV tat, and also a region activated by the endogenous **nuclear**
factor NF-.**kappa**.**B** (the LTR has tandem NF-.kappa.B binding regions). Although the tar sequence strongly suppresses expression in the absence of tat (see. . .

DETDESC:

DETD(7)

The . . . folding enzymes, transport proteins, and the like), down-regulating host cell regulatory factors employed by the infectious agent (for example, the NF-.**kappa**.**B** **nuclear** **factor** found in activated lymphocytes which up-regulates HIV-1 transcription), up-regulating viral or host cell factors which suppress viral gene expression, by. . .

US PAT NO: 5,854,223 [IMAGE AVAILABLE] DATE ISSUED: Dec. 29, 1998

L15: 10 of 72

S-DC28 as an antirestenosis agent after balloon injury

INVENTOR:

OR: Cy Stein, New City, NY
LeRoy Rabbani, New York, NY
E: The Trustees of Columbia University in the City of New

ASSIGNEE: York, New York, NY (U.S. corp.)

APPL-NO: 08/678,234

DATE FILED: Jul. 11, 1996

PRIM-EXMR: Scott W. Houtteman

LEGAL-REP: John P.Cooper & Dunham LLP White

US PAT NO: 5,854,223 [IMAGE AVAILABLE] L15: 10 of 72

US-CL-CURRENT: **514/44**; 536/24.5

DETDESC:

DETD(109)

In . . . non-sequence specifically inhibit fibronectin's binding to

US PAT NO: 5,849,286 [IMAGE AVAILABLE] DATE ISSUED: Dec. 15, 1998

L15: 11 of 72

TTTLE: Ubiquitin conjugating enzymes 7,8 and 9
INVENTOR: Jian Ni, Gaithersburg, MD
Reiner Gentz, Silver Springs, MD

Mark D. Adams, North Potomac, MD

Human Genome Sciences, Inc., Gaithersburg, MA (U.S. corp.) 08/464,604 ASSIGNEE:

APPL-NO: 08/464,604 DATE FILED: Jun. 5, 1995

PRIM-EXMR: Pot Robert A. Wax Lisa J. Hobbs

Elliot M. Olstein, J. G. Mullins LEGAL-REP:

L15: 11 of 72

US PAT NO: 5,849,286 [IMAGE AVAILABLE] US-CL-CURRENT: 424/94.5; 435/193; **514/12*

BSUM(10)

Maturation of the p105 **NF**-**KB** precursor into the active p50 subunit of the transcriptional activator also proceeds in a ubiquitin and proteasome-dependent manner. Furthermore, inhibitors to the proteasome block degradation of IkBa and thus prevent tumor necrosis factor alpha induced activation of **NF**-**KB** and its entry into the nucleus.

US PAT NO: 5,849,263 [IMAGE AVAILABLE] DATE ISSUED: Dec. 15, 1998

Pharmaceutical compositions containing alkylaryl polyether

alcohol polymer

Thomas P. Kennedy, Richmond, VA INVENTOR:

Charlotte-Mecklenburg Hospital Authority, Charlotte, NC ASSIGNEE:

(U.S. corp.)

APPL-NO: 08/638,893 DATE FILED: Apr. 25, 1996 ART-UNIT: 166

PRIM-EXMR: Robert H. Harrison

LEGAL-REP: Bell Seltzer Intellectual Property Law Group of Alston &

Bird LLP

US PAT NO: 5,849,263 [IMAGE AVAILABLE] L15: 12 of 72 US-CL-CURRENT: 424/45, 78.05, 78.06, 78.08, 78.37; **514/179**, **885**, **887**

SUMMARY:

BSUM(14)

These cytokines share regulation of their expression by the transcription factor **Nuclear** **Factor** **kappa**.**B** (NF-.**kappa**.B), a particularly important transcription factor mediating inflammatory events (U. Siebenlist, G. Granzuso and R. Brown. "Structure, regulation and function of. . . human peripheral blood mononuclear cells". International Journal of Immunology (1993) 6:409-422; mononuclear cells. International Journal of Immunology (1993) 6:409

**R. Schreck, et al. "Dithiocarbamates as potent inhibitors of **nuclear*

factor.**kappa**.**B** activation in intact cells*. Journal of

Experimental Medicine (1992) 175:1181-1194). However, the few

antioxidants known to inhibit NF-.kappa.B activation share.

US PAT NO: 5,846,961 [IMAGE AVAILABLE] DATE ISSUED: Dec. 8, 1998 L15: 13 of 72

Multi-faceted method to repress reproduction of latent TITLE:

viruses in humans and animals

Knox Van Dyke, Morgantown, WV
HIV Diagnostics, Inc., Lexington, KY (U.S. corp.) INVENTOR: ASSIGNEE:

APPL-NO: 08/479,010

DATE FILED: Jun. 7, 1995 152

ART-UNIT: PRIM-EXMR:

Gollamudi S. Kishore LEGAL-REP:

Price, Heneveld, Cooper, DeWitt & Litton

US PAT NO: 5,846,961 [IMAGE AVAILABLE] L15: 13 of 72 US-CL-CURRENT: **514/171**, **198**, **369**, **374**, **378**, **561**.

ABSTRACT:

Disclosed . . . such as HIV, in animals by the generally concurrent administration of (1) antioxidants including a glutathione agent; and (2) an **NFKB** induction inhibitor. Also disclosed are pharmaceutical

compositions and kits for use in repressing reproduction of latent viruses such as HIV.

SUMMARY:

BSUM(8)

Schreck . . . The iKB factor is removed from the protein triad and the remaining p50, p65 complex becomes known as NF-kappa B (**NFKB**).

SUMMARY:

BSUM(9)

Schreck et al. have recognized that **NFKB** is a gene transcription factor that migrates into the nucleus of the HIV infected cell and switches on the production. . . expression of HIV-I in a human T cell line. They further report that the expression of HIV is mediated by **NFKB** transcription factor which is potently and rapidly activated by a hydrogen peroxide treatment of cells from its inactive cytoplasmic form. They additionally report that N-acetyl cysteine and other thiol compounds block the activation of **NFKB**. They concluded that these diverse agents thought to activate **NFKB** by distinct intracellular pathways might act through a common mechanism involving the synthesis of reactive oxygen intermediates. They did not. . .

SUMMARY:

BSUM(10)

Sherman et al., Biochem. Biophys. Res. Comm., 191 (3):1301-1308, 1993, report that pyrrolidine dithiocarbamate (PDTC) is an inhibitor of "*NFKB** activation. They further report that this compound is an inhibitor of nitric oxide synthase (NO synthase). They further report that. . . that PDTC may act as a scavenger of reactive oxygen species which prevents them from participation in the activation of **NFKB**.

SUMMARY:

BSUM(15)

The . . . the generally concurrent administration of 1) a glutathione agent; 2) at least one additional antioxidant; and 3) at least one "NFKB*" induction inhibitor. Further aspects and advantages of the invention will be apparent to those skilled in the art upon review. . .

DETDESC:

DETD(3)

There . . . glutathione precursor, a glutathione production enhancer, or glutathione, (2) high doses of additional fat- and water-soluble antioxidants, and (3) an **NFKB** induction inhibitor, to an animal infected with a latent virus. The fat- and water-soluble antioxidants are administered to an animal. . .

DETDESC:

DETD(5)

The Role of **NFKB** and Peroxynitrite in the Activation of a Cell to Reproduce HIV **NFKB** is a gene transcription factor that switches on the production of the HIV virus of a virally infected cell. **NFKB** is known to activate a variety of genes, including the transcription of a variety of cytokines, viruses and NO Synthase.. . .

DETDESC:

DETD(9)

Peroxynitrite is significant in that it activates **NFKB**. **NFKB** is inactivated by I Kappa B (IKB) which acts on **NFKB** via the P65 subunit. As shown in FIG. 1, peroxynitrite cleaves IKB, thereby releasing the active **NFKB**.

DETDESC:

DETD(31)

NFKB Induction Inhibitors

DETDESC:

DETD(32)

NFKB induction inhibitors are agents that inhibit **NFKB** transcription factor from binding to DNA. This blocks the induction of HIV or other viral reproduction by directly suppressing the viral reproduction activating mechanism. **NFKB** inhibitors (item 7, FIG. 2) also suppress peroxynitrite synthesis, by preventing **NFKB** from

activating cell genes to produce NO synthase.

DETDESC:

DETD(35)

The preferred type of **NFKB** induction inhibitor is an anti-inflammatory steroid. Examples of suitable anti-inflammatory steroids suitable as **NFKB** induction inhibitors include but are not limited to predonsone, prednisolone, methyl prednisolone, dexamethasone, beta metasone dehydroepiandrosterone, 9a-fluorocortisol, prednisone, aetiocholanolone, 2-methylcortisol, pregnanediol, deoxycorticosterone, cortisone, hydrocortisone (cortisol), 6a-methylprednisolone, triamcinolone, estrogen or derivatives thereof. Generally, any steroid with antiinflammatory action toward **NFKB** may be used. In addition, one or more suitable nonglucocorticoid lazaroids may be utilized as **NFKB** induction inhibitors. Preferred lazaroids include, but are not limited to, U-74006F, which is 21-[4-(2,6-di-1-pyrrolidinyl-4-pyrimidinyl)-1-piperazily] -1-6-methyl-(16alpha,-pregna-1,4.9(11)triene-3,20-dione monomethanesulfonate or TIRILAZAD mesylate or. . .

DETDESC:

DETD(39)

In . . . weight. Other antiinflammatory steroids can be substituted at appropriate doses, as set forth in the Physicians' Desk Reference. Adminstration of an **NFKB** induction inhibitor such as an anti-inflammatory steroid, is one of the most important steps in the treatment of HIV, AIDS. . .

DETDESC:

DETD(43)

In addition, to the previously noted anti-inflammatory steroids and lazaroids, a variety of other compounds may be utilized as "NFKB" induction inhibitors such as pyrrolidine dithiocarbamate and other dithiocarbamates, and glycyrrhizic acid (from licorice root). A preferred dosage level when. . . is about 100 mg/day per person for each day of therapy. In addition, other compounds are suitable for use as ""NFKB" induction inhibitors. These inhibitors include, but are not limited to, immunosuppressants such as cyclosporin A, rapamycin, interleukin 10, and FK. . . Clearly, a wide array of plant steroids, male steroids, female steroids, glucocorticoids, lazaroids, and 21-aminosteroids are eligible for use as "NFKB" induction inhibitors.

DETDESC:

DETD(44)

An inhibitor known to be effective against **NFKB** binding or expressing is mevinolin, a drug which prevents isoprenylation and methylthioadenosine (MTA) and inhibitor of several S adenosylmethionine dependent. . .

DETDESC:

DETD(51)

Although . . . antioxidants, glutathione agents, and steroids with regard to HIV production. HIV replication is blocked by a combination of antioxidants and **NFKB** induction inhibitor. About 7% of the blocking action of HIV replication is believed to stem from the **NFKB** induction inhibitor, which preferably is one or more anti-inflammatory steroids. Although such steroids do not have direct inhibitory activity, they control viral synthesis by blocking **NFKB** induction. As will be recalled, **NFKB** is a DNA transcription factor made of protein. **NFKB** controls a whole series of inflammatory cytokines and NO synthase as well as HIV and FIV replication. Upon introduction of. . .

DETDESC:

DETD(52)

However, for ""NFKB"" to be active it must shed its inhibitory factor I kappa B. Such shedding requires oxidation because the bonds holding. . to proteins P50 and P65 are sensitive to oxidation. Thus, antioxidants keep the I kappa B inhibitory factor bound to ""NFKB" and therefore inactive. The role of antioxidants in the mechanism depicted in FIG. 3 is believed to be responsible for about 30% of the activity of producing ""NFKB" and preventing HIV replication.

DETDESC:

DETD(53)

All... known to those skilled in the art. Although it is most preferred to administer the anitoxidants including glutathione agent and **NFKB** induction inhibitor concurrently, or simultaneously, it is not a

requirement. Thus, the preferred embodiments of the present invention

DETDESC:

DETD(55)

The . . . a glutathione agent; (2) an effective amount of one or more additional antioxidants; and (3) an effective amount of an **NFKB** induction inhibitor. In a most preferred embodiment, the pharmaceutical compositions comprise; (1) an effective amount of a glutathione agent, e.g. . . antioxidant, (2b) an effective amount of a fat-soluble antioxidant, and (3) an effective amount of an anti-inflammatory steroid as the **NFKB** induction inhibitor. The other ingredients described above may also be included.

DETDESC:

DETD(62)

In . . . C, A and E; an effective amount of at least one glutathione precursor such as N-acetyl cysteine; followed by an "*NFKB** induction inhibitor such as one or more anti-inflammatory steroids or lazaroids. As summarized in Table 4 below, seven cats heavily. . . 10 to about 18 pounds. The cats were initially treated with a single dosage of an effective amount of an "*NFKB** induction inhibitor, that is an antiinflammatory steroid dose of DEPO-MEDROL (20-25 mg) and a series of oral dosages of a. . .

DETDESC:

DETD(66)

In . . . fat-soluble antioxidants and an effective amount of at least one glutathione precursor such as N-acetyl cysteine are administered. Before an **NFKB** induction inhibitor is administered, the CD.sub.4 (T-lymphocyte) count is increased to about 100 cells/mm.sup.3 or more. The CD.sub.4 count may. . . concentrates containing monocytes may be given, such as via transfusions. Once CD.sub.4 counts are about 100 cells/mm.sup.3 or more, an **NFKB** induction inhibitor is administered.

DETDESC:

DETD(67)

In both the preferred and optional treatment regimens, the **NFKB** induction inhibitor is administered until AIDS(-) is indicated from AIDS(+) blood assay, via ELISA, Western blot, and PCR (polymerase chain.

DETDESC:

DETD(78)

Preferably, . . . suitable glutathione precursors could be utilized in place of, or instead of the N-acetyl cysteine. Similarly, one or more other **NFKB** induction inhibitors could be utilized in place of or instead of the methyl prednisolone.

DETDESC:

DETD(81)

The . . . one fat soluble antioxidant at doses higher than the recommended daily minimum requirements, and preferably, only slight amounts or no "NFKB" induction inhibitor. In a most preferred treatment regimen, the subject suffering from symptoms of the Herpes virus is administered generally. . .

CLAIMS:

CLMS(1)

The .

combinations thereof,
ii) at least one additional antioxidant at doses higher than the
recommended daily minimum requirements; and
iii) at least one **NFKB** induction inhibitor in an amount effective to

iii) at least one ""NFKB" induction inhibitor in an amount effective to inhibit "nuclear" "factor" "kappa" "B"; said at least one "NFKB" induction inhibitor being selected from the group consisting of anti-inflammatory steroids and nonglucocorticoid lazaroids.

CLAIMS:

CLMS(11)

11. The method of claim 1 wherein said **NFKB** induction inhibitor comprises at least one anti-inflammatory steroid.

CLAIMS:

CLMS(13)

13. The method of claim 1 wherein said **NFKB** induction inhibitor comprises a nonglucocorticoid lazaroid.

CLAIMS:

CLMS(15)

The method of claim 1 further comprising administering:
 a peroxynitrite production suppressor in addition to said **NFKB** induction inhibitor.

CLAIMS:

CLMS(17)

17. . . .

combinations thereof:

ii) at least one additional antioxidant at doses higher than the recommended daily minimum requirements; and iii) at least one **NFKB** induction inhibitor in an amount effective to

inhibit ""nuclear" ""factor" ""kappa" ""B""; said at least one
""NFKB" induction inhibitor being selected from the group consisting
of anti-inflammatory steroids and nonglucocorticoid lazaroids.

CLAIMS:

CLMS(27)

27. The method of claim 17 wherein said **NFKB** induction inhibitor comprises at least one anti-inflammatory steroid.

CLAIMS:

CLMS(29)

29. The method of claim 17 wherein said **NFKB** induction inhibitor comprises a nonglucocorticoid lazaroid.

CLAIMS:

CLMS(31)

The method of claim 17 further comprising administering:
 iv) a peroxynitrite production suppressor in addition to said **NFKB** induction inhibitor.

US PAT NO: 5,846,959 [IMAGE AVAILABLE]

L15: 14 of 72

DATE ISSUED: Dec. 8, 1998

TITLE: Treatment for atherosclerosis and other cardiovascular and

inflammatory diseases

INVENTOR: Russell M. Medford, Atlanta, GA

R. Wayne Alexander, Atlanta, GA Sampath Parthasarathy, Atlanta, GA Bobby V. Khan, Dunwoody, GA

ASSIGNEE: Emory University, Atlanta, GA (U.S. corp.)

APPL-NO: 08/471,537

DATE FILED: Jun. 6, 1995

ART-UNIT: 161

PRIM-EXMR: Peter O'Sullivan

LEGAL-REP: Sherry M. Knowles, JacquelineKing & Spalding Haley

US PAT NO: 5,846,959 [IMAGE AVAILABLE] L15: 14 of 72 US-CL-CURRENT: **514/165**; 424/9.1, 9.2; 435/6, 7.2, 7.21, 7.24, 7.94, 7.95; 436/71, 86, 129, 172, 503, 504, 548; **514/18**, **171**, **211**, **423**, **457**, **478**, **479**

SUMMARY:

BSUM(5)

Molecular . . . of the regulatory elements on the human VCAM-1 gene that control its expression suggests an important role for nuclear factor-kB (**NF**-*KB**), a transcriptional regulatory factor, or an NF-k-beta. like binding protein in oxidation-reduction-sensitive regulation of VCAM-1 gene expression. Transcriptional factors are. . role in mediating inflammatory and other stress signals to the nuclear regulatory apparatus. Although the precise biochemical signals that activate **NF**-**kB** are unknown, this transcriptional factor may integrate into a common molecular pathway many of the risk factors and *causative* signals.

SUMMARY:

BSUM(6)

Importantly, the activation of **NF**-**kB** in vascular endothelial

cells by diverse signals can be specifically inhibited by antioxidants such as N-acetyleysteine and pyrrolidine dithiocarbamate (see. 0.70969,934, now allowed). This has led to the hypothesis that oxygen radicals play an important role in the activation of "*NF"---*RB"* through an undefined oxidation-reduction mechanism. Because an "*NF"---*Ike enhancer element also regulates the transcription of the VCAM-1 promoter in an oxidation-reduction-sensitive manner, oxidative stress in the atherosclerotic lesion. . .

DRAWING DESC:

DRWD(8)

FIG. . . . an illustration of an autoradiogram that indicates that linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive **NF**-**kB** like factor. RAEC were split at the ratio to give approximately 60% confluence in 100-mm tissue culture plates. HAEC were. . .

DRAWING DESC:

DRWD(9)

FIG. 8 is an illustration of an acrylamide gel slab that indicates that polyunsaturated fatty acids activate **NF**.***RB**-like DNA binding activities that are blocked by the antioxidan PDTC. Confluent HAEC in media containing 4% FBS (as described in. . . native acrylamide gels, and exposed to autoradiography film at -70.degree. C. for 18 hours. Two bands A and C, representing **NF**-**kB*** like binding activity are designated. A weak band B was observed in control (untreated) cells.

DETDESC:

DETD(31)

Previous . . . promoter studies that cytokines and non-cytokines activate VCAM-1 gene expression in endothelial cells at least in part transcriptionally through two **NF*-***RB***.like DNA binding elements. It has also been demonstrated that PDTC inhibits VCAM-1 gene expression through a redox-sensitive **NF*-**RB***.like factor. To determine whether polyunsaturated fatty acids induce transcriptional activation of the human VCAM-1 promoter via a similar mechanism, . . . results were obtained with the minimal cytokine-inducible promoter of the VCAM-1 gene (p85 VCAM-CAT), containing the -77 and -63 bp **NF*-***RB**-like sites. Neither linoleic acid nor TNF-alpha. had any effect on activity using a constitutively expressed pSV.sub.2 CAT construct. PDTC inhibited . . indicate that analogous to TNF-alpha. polyunsaturated fatty acids such as linoleic acid induce the transcriptional activation of VCAM-1 through an **NF*-**RB**-like redox-sensitive mechanism.

DETDESC:

DETD(32)

To determine whether polyunsaturated fatty acids and their oxidative metabolites regulate VCAM-1 promoter activity through an **NF**.**kB**-like transcriptional regulatory factor, nuclear extracts from HAEC were assayed for DNA binding activity to a double-stranded oligonucleotide containing the VCAM-1 **NF**.**kB**-like promoter elements located at positions -77 and -63. As shown in FIG. 7, two bands A and C, representing **NF**.**kB**-like activity were induced in response to a three hour exposure to linoleic acid (7.5 .mu.M). Similar findings were observed on. . . exposure to the cytokine TNF-.alpha. (100 U/ml). A weak band B was observed in control (untreated) cells. No induction of **NF**.**kB**-like binding was observed with the monounsaturated fatty acid oleic acid. Pretreatment of the cells for thirty minutes with PDTC inhibited. . . previously reported findings that PDTC blocks the activation of VCAM-1 gene expression in HUVEc by inhibiting the activation of these **NF*****KB**-like DNA binding proteins.

DETDESC:

DETD(55)

Linoleic Acid Induces Transcriptional Activation of the VCAM-1 Promoter by a Redox-Sensitive **NF**-**kB** Like Factor

DETDESC:

DETD(57)

FIG. 7 illustrates the results of this experiment. Linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive "NF"-1 like factor. These results are similar to those observed by the activation of VCAM-1 promotor by cytokines such as TNF-alpha...

DETDESC:

DETD(59)

Polyunsaturated Fatty Acids Activate **NF**-**kB**-like DNA Binding Activities that are Blocked by the Antioxidant PDTC

DETDESC

DETD(60)

Confluent... native acrylamide gels, and exposed to autoradiography film at -70.degree. C. for 18 hours. Two bands A and C, representing ""NF"--"kB" like binding activity are designated. A weak band B was observed in control (untreated) cells.

US PAT NO: 5,843,643 [IMAGE AVAILABLE] L15: 15 of 72

DATE ISSUED: Dec. 1, 1998

TITLE: Site-specific transfection of eukaryotic cells using polypeptide-linked recombinant nucleic acid

INVENTOR: Paul L. Ratner, 11 Ash St., Bar Harbor, ME 04609

APPL-NO: 08/199,608

DATE FILED: Feb. 22, 1994

ART-UNIT: 187

PRIM-EXMR: W. Gary Jones

ASST-EXMR: Dianne Rees

LEGAL-REP: Patrick D. Kelly

US PAT NO: 5,843,643 [IMAGE AVAILABLE] L15: 15 of 72 US-CL-CURRENT: 435/6, 5, 91.1; =*514/2=*, **44**; 530/300, 350; 536/23.1, 24.3, 24.5

DETDESC:

DETD(12)

. . .

SRF c-fos oncogene

Thyroid hormone Growth hormone

receptor

CREB Somatostatin Singh et al 1989
NF-**kB** Human immunodeficiency virus

YB-1 Major histocompatibility II

GATA-1 Beta globin Trainor et al 1990

GCF Epidermal. . .

DETDESC:

DETD(246)

linear

(ii) MOLECULE TYPE: DNA (genomic)

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO
(vi) ORIGINAL SOURCE:

(A) ORGANISM: binding site for **NF**-**KB** chromosome binding

protein

(B) STRAIN: human

(F) TISSUE TYPE: human (G) CELL TYPE: human

(x) PUBLICATION INFORMATION:

(A) AUTHORS: Singh,. .

US PAT NO: 5,840,710 [IMAGE AVAILABLE] L15: 16 of 72

DATE ISSUED: Nov. 24, 1998

TITLE: Cationic amphiphiles containing ester or ether-linked

lipophilic groups for intracellular delivery of

therapeutic molecules INVENTOR: Edward R. Lee,

OR: Edward R. Lee, Quincy, MA David J. Harris, Lexington, MA Craig S. Siegel, Woburn, MA

Mathieu B. Lane, Cambridge, MA Shirley C. Hubbard, Belmont, MA Seng H. Cheng, Wellesley, MA

Simon J. Eastman, Marlboro, MA John Marshall, Milford, MA

Ronald K. Scheule, Hopkinton, MA
ASSIGNEE: Genzyme Corporation, Framingham, MA (U.S. corp.)

APPL-NO: 08/546,087 DATE FILED: Oct. 20, 1995 ART-UNIT: 189

ART-UNIT: 189
PRIM-EXMR: Bruce R. Campbell

LEGAL-REP: Finnegan, Henderson, Farabow, Garrett & Dunner, L.L.P.

US PAT NO: 5,840,710 [IMAGE AVAILABLE] L15: 16 of 72

US-CL-CURRENT: **514/44**; 424/450; **514/2**; 554/1, 227; 560/1, 224

DETDESC:

DETD(285)

It . . . increase with the severity of an inflammatory condition (for example, tumor necrosis factor "TNF" and potentially transcription factors such as **NF**-**kB**, AP-1, NF-IL6 and octamer binding protein). It has also been determined that interleukin 8, a polypeptide of 8,500

US PAT NO: 5,840,277 [IMAGE AVAILABLE] DATE ISSUED: Nov. 24, 1998 L15: 17 of 72 TITLE: Treatment of chronic pulmonary inflammation INVENTOR: Andrew J. Ghio, Durham, NC Thomas P. Kennedy, Richmond, VA ASSIGNEE: Charlotte Hospital Authority, Charlotte, NC (U.S. corp.) 08/632,275 APPL-NO: DATE FILED: Apr. 15, 1996 185 ART-UNIT: PRIM-EXMR: David Guzo LEGAL-REP: The Bell Seltzer Intellectual Law Firm of Alston & Bird, LLP

US PAT NO: 5.840.277 [IMAGE AVAILABLE] L15: 17 of 72 US-CL-CURRENT: 424/45, 78.05, 78.37; **514/828**, **851**

SUMMARY:

BSUM(3)

The . . . inflammation. More particularly, the present invention relates to the use of alkylaryl polyether alcohol polymers to reduce the activation of **nuclear****factor****kappa****B** (NF-.**kappa**.B) and inhibit the secretion of pro-inflammatory cytokines TNF-alpha (TNF-.alpha.), interleukin-1 beta (IL-1.beta.), interleukin-6 (IL-6), interleukin-8 (IL-8) and the growth factor. .

SUMMARY:

BSUM(15)

These cytokines share regulation of their expression by the transcription factor **Nuclear** **Factor** **kappa**-**B** (NF-.**kappa**.B), a particularly important transcription factor mediating inflammatory events (U. Siebenlist, G. Granzuso and R. Brown. "Structure, regulation and function of. . . human peripheral blood mononuclear cells". International Journal of Immunology (1993) 6:409-422; R. Schreck, et al. *Dithiocarbamates as potent inhibitors of **nuclear* **factor** .**kappa** .**B** activation in intact cells*. Journal of Experimental Medicine (1992) 175:1181-1194). However, the few antioxidants known to inhibit NF- kappa.B activation share. . .

US PAT NO: 5,837,510 [IMAGE AVAILABLE]
DATE ISSUED: Nov. 17, 1998 1.15: 18 of 72 Methods and polynucleotide constructs for treating host

cells for infection or hyperproliferative disorders
R: Mark A. Goldsmith, 263 Chenery St., San Francisco, CA INVENTOR: 94131

Robert O. Ralston, 2863 Judah, San Francisco, CA 94122

APPL-NO: 08/472,056 DATE FILED: Jun. 6, 1995 ART-UNIT: 185 PRIM-EXMR: Johnny Railey

LEGAL-REP: Norman J. Kruse, Donald J. Pochopien, Robert P. Blackburn

5,837,510 [IMAGE AVAILABLE] US-CL-CURRENT: 435/455; 424/93.2; 435/320.1, 456; **514/44**; 536/23.1, 23.2, 23.5, 23.53, 23.6, 23.7, 23.72, 24.1, 24.5

DETDESC:

DETD(6)

The . . . contains both the tar region, which is highly selective for HIV tat, and also a region activated by the endogenous **nuclear**

factor NF..sub..**kappa**. **B** (the LTR has tandem NF..sub..kappa.

B binding regions). Although the tar sequence strongly suppresses expression in the absence of tat. .

DETDESC:

DETD(7)

The . . . folding enzymes, transport proteins, and the like), down-regulating host cell regulatory factors employed by the infectious agent (for example, -the NF-.sub..**kappa**. **B** **nuclear** **factor** found in activated lymphocytes which up-regulates HIV-1 transcription), up-regulating viral or host cell factors which suppress viral gene expression, by. . .

US PAT NO: 5,830,848 [IMAGE AVAILABLE] DATE ISSUED: Nov. 3, 1998

Method and agents for inducement of endogenous nitric oxide synthase for control and management of labor TITLE: during pregnancy

OR: Michael R. Harrison, San Francisco, CA Michael A. Heymann, San Francisco, CA INVENTOR: Robert Kirk Riemer, Half Moon Bay, CA Eileen Stack Natuzzi, San Francisco, CA

ASSIGNEE: The Regents of the University of California, Oakland, CA (U.S. corp.)

APPL-NO: 08/450,126 DATE FILED: May 25, 1995 ART-UNIT: PRIM-EXMR: 182 Stephen Walsh ASST-EXMR: Daryl A. Basham LEGAL-REP: Hana Verny

5,830,848 [IMAGE AVAILABLE] US-CL-CURRENT: **514/2**; 424/85.1, 85.2, 85.5; 530/399

DETDESC:

DETD(25)

Additionally, . . . on NO production augmentation. These agents are putative control elements which modify the expression of transcriptional regulatory proteins such as ""nuclear" "Tactor" NF "Kappa" "B"

Jun/fos, tumor necrosis factor (TNT-alpha.), NF-116, activator protein (AP-1), octamer binding protein, (OCT-1), (OCT-2), PU-1, and gamma activation factor (GAF),. . .

CLAIMS:

CLMS(19)

19. The method of claim 1 wherein the transcriptional regulating protein is **nuclear** **factor** **kappa** **B** Jun/fos.

US PAT NO: 5,824,664 [IMAGE AVAILABLE] L15: 20 of 72 DATE ISSUED: Oct. 20, 1998

TITLE: Suppression of HIV expression by organic thiophosphate INVENTOR: Philip S. Schein, Bryn Mawr, PA

Thea Kalebic, Bethesda, MD

ASSIGNEE: U.S. Bioscience, Inc., West Conshohocken, PA (U.S. corp.)

National Institutes of Health, The National Cancer
Institute, Rockville, MD (U.S. corp.)

APPL-NO: 08/037,633

DATE FILED: Mar. 26, 1993

ART-UNIT: 188

PRIM-EXMR: Leon B. Lankford, Jr. Francisco C. Prats ASST-EXMR:

LEGAL-REP: Pennie & Edmonds LLP

US PAT NO: 5,824,664 [IMAGE AVAILABLE] US-CL-CURRENT: **514/143**, **75**, **114**, **665** L15: 20 of 72

DETDESC:

DETD(73)

Similar to N-acetyl cysteine, which suppressed PMA- and TNF alpha -mediated induction of HIV-LTR transcription by inhibiting **NFKB** activity (Staal, F. J. et al., 1990, supra), WR 151327 suppressed transcriptional activity of HIV-LTR in transiently transfected RD cells.. . .

US PAT NO: 5,821,260 [IMAGE AVAILABLE] DATE ISSUED: Oct. 13, 1998

Treatment for atherosclerosis and other cardiovascular and

L15: 21 of 72

inflammatory diseases
INVENTOR: Russell M. Medford, Atlanta, GA
Margaret K. Offermann, Atlanta, GA

R. Wayne Alexander, Atlanta, GA Sampath Parthasarathy, Atlanta, GA

ASSIGNEE: Emory University, Atlanta, GA (U.S. corp.) 08/485,307

APPL-NO: DATE FILED: Jun. 7, 1995 ART-UNIT: 129

PRIM-EXMR: Peter O'Sullivan

LEGAL-REP: Sherry M. Knowles, JacquelineKing & Spalding Haley

US PAT NO: 5,821,260 [IMAGE AVAILABLE] L15: 21 of 72

```
US-CL-CURRENT: **514/423**; 424/9.1, 9.2; 435/6, 7.2, 7.21, 7.24, 7.94, 7.95; 436/71, 86, 129, 172, 503, 504, 548; **514/18**, **226.2**, **477**, **478**, **479**, **484**, **485**, **478**, **484**, **485**, **487**
```

DETDESC:

DETD(63)

At the molecular level, PDTC has been shown to inhibit the activation of the transcriptional regulatory factor **Nf*--*kB** in response to certain cytokine and non-cytokine stimuli (Schreck, Rieber et al. 1991; Schreck, Meier et al. 1992). However, by. . . has been discovered that endothelial cells activate VCAM-1 gene expression through an apparently novel transcriptional regulatory factor that is not **Nf*--*kB**. This suggests that PDTC may regulate endothelial cell gene expression through its effect on a new transcriptional regulatory protein. It. . . .

```
US PAT NO: 5,814,612 [IMAGE AVAILABLE]
DATE ISSUED: Sep. 29, 1998
                                                                               L15: 22 of 72
                 Retinol derivatives and uses thereof
INVENTOR:
                     Jochen Buck, New York, NY
              Ulrich Hammerling, New York, NY
Fadila Derguini, New York, NY
              Koji Nakanishi, New York, NY
             E: Sloan-Kettering Institute for Cancer Research, New York, NY (U.S. corp.)
The Trustees of Columbia in the City of New York, New
ASSIGNEE:
                York, NY (U.S. corp.)
 APPL-NO:
                    07/880,041
DATE FILED: May 6, 1992
ART-UNIT:
PRIM-EXMR:
                      Johann Richter
                     John Peabody
John P. White
ASST-EXMR:
LEGAL-REP:
US PAT NO: 5,814,612 [IMAGE AVAILABLE] L15: 22 of 72 US-CL-CURRENT: **514/21**, **725**; 549/453, 512, 551, 554, 561, 563; 552/10, 11, 12; 558/430; 562/867; **564/123**, **152**,
                **153**; 568/823, 824, 825
```

DETDESC:

DETD(81)

The . . . molecules, that shuttle to the nucleus to regulate transcription. For example, the protein products of the rel gene family (e.g., "*NF*--**kB**) translocate upon activation from the cytoplasm to the nucleus and regulate transcription (24). Small lipophilic molecules including the steroids, vitamin. . .

```
US PAT NO: 5,811,449 [IMAGE AVAILABLE]
DATE ISSUED: Sep. 22, 1998
                                                                                   L15: 23 of 72
                   Treatment for atherosclerosis and other cardiovascular and
TITLE:
                 inflammatory diseases
              DR: Russell M. Medford, Atlanta, GA
R. Wayne Alexander, Atlanta, GA
INVENTOR:
               Sampath Parthasarathy, Atlanta, GA
              Bobby V. Khan, Dunwoody, GA
                    Emory University, Altanta, GA (U.S. corp.) 08/483,335
ASSIGNEE:
APPL-NO:
DATE FILED: Jun. 7, 1995
ART-UNIT:
                     129
PRIM-EXMR:
                      Peter O'Sullivan
                      Sherry M. Knowles, JacquelineKing & Spalding Haley
LEGAL-REP:
US PAT NO: 5,811,449 [IMAGE AVAILABLE]
US-PAT-NO: 5,811,449 [IMAGE AVAILABLE] L15: 23 of 7
US-CL-CURRENT: **514/423**; 424/9.1, 9.2; 436/71, 86, 129; **514/
**226,2**, **477**, **478**, **479**, **484**, **485**,
**487**, **488**, **489**, **506**, **513**, **517**,
**518**, **553**, **561**, **824**, **825**, **826**,
**861**, **863**; 530/331; 548/431, 531; 549/10;
                558/230, 234, 235, 250; 562/26, 27; **564/76**; 568/21,
```

SUMMARY:

BSUM(5)

Molecular . . . of the regulatory elements on the human VCAM-1 gene that control its expression suggests an important role for nuclear factor-kB (**NF**-*kB**), a transcriptional regulatory factor, or an NF-k.beta. like binding protein in oxidation-reduction-sensitive regulation of VCAM-1 gene expression. Transcriptional factors are. . role in mediating inflammatory and other stress signals to the nuclear regulatory apparatus. Although the precise biochemical signals that activate **NF**-**kB** are unknown, this transcriptional factor may integrate into a common molecular pathway many of the risk factors and

"causative" signals. . .

SUMMARY:

BSUM(6)

Importantly, the activation of ""NF"-""kB" in vascular endothelial cells by diverse signals can be specifically hinhibited by antioxidants such as N-acetyleysteine and pyrrolidine dithiocarbamate (see. 07/969,934, now allowed). This has led to the hypothesis that oxygen radicals play an important role in the activation of ""NF"-""kB" through an undefined oxidation-reduction mechanism. Because an ""NF"-""kB"-like enhancer element also regulates the transcription of the VCAM-I promoter in an oxidation-reduction-sensitive manner, oxidative stress in the atherosclerotic lesion.

DRAWING DESC:

DRWD(8)

FIG. . . . an illustration of an autoradiogram that indicates that linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive **NF**-**kB** like factor. HAEC were split at the ratio to give approximately 60% confluence in 100-mm tissue culture plates. HAEC were. . .

DRAWING DESC:

DRWD(9)

FIG. 8 is an illustration of an acrylamide gel slab that indicates that polyunsaturated fatty acids activate "*NF"---"*RB"--like DNA binding activities that are blocked by the antioxidant PDTC. Confluent HAEC in media containing 4% FBS (as described in. . . native acrylamide gels, and exposed to autoradiography film at -70.degree. C. for 18 hours. Two bands A and C, representing **NF"*--*kB"*-like binding activity are designated. A weak band B was observed in control (untreated) cells.

DETDESC:

DETD(31)

Previous . . . promoter studies that cytokines and non-cytokines activate VCAM-1 gene expression in endothelial cells at least in part transcriptionally through two **NF*-*******NB***.like DNA binding elements. It has also been demonstrated that PDTC inhibits VCAM-1 gene expression through a redox-sensitive **NF*-***LB***.like factor. To determine whether polyunsaturated fatty acids induce transcriptional activation of the human VCAM-1 promoter via a similar mechanism, . . . results were obtained with the minimal cytokine-inducible promoter of the VCAM-1 gene (p85 VCAM-CAT), containing the -77 and -63 bp **NF**-***RB**-like sites. Neither linoleic acid nor TNF-alpha. had any effect on activity using a constitutively expressed pSV.sub.2 CAT construct. PDTC inhibited. . . indicate that analogous to TNF-alpha. polyunsaturated fatty acids such as linoleic acid induce the transcriptional activation of VCAM-1 through an **NF**-***RB**-like redox-sensitive mechanism.

DETDESC:

DETD(32)

To determine whether polyunsaturated fatty acids and their oxidative metabolites regulate VCAM-1 promoter activity through an "NF"*-"*KB*-like transcriptional regulatory factor, nuclear extracts from HAEC were assayed for DNA binding activity to a double-stranded oligonucleotide containing the VCAM-1 ""NF"*-"*kB*-like promoter elements located at positions -77 and -63. As shown in FIG. 7, two bands A and C, representing "*NF"*-"*kB*-like activity were induced in response to a three hour exposure to linoleic acid (7.5 .mu.M). Similar findings were observed on. . . exposure to the cytokine TNF-alpha. (100 U/ml). A weak band B was observed in control (untreated) cells. No induction of "*NF"*-"*kB*-like binding was observed with the monounsaturated fatty acid oleic acid. Pretreatment of the cells for thirty minutes with PDTC inhibited. . . previously reported findings that PDTC blocks the activation of VCAM-1 gene expression in HUVEC by inhibiting the activation of these "*NF"*- "*KB"*-like DNA binding proteins.

DETDESC:

DETD(55)

Linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive **NF**-**kB** like factor

DETDESC:

DETD(57)

FIG. 7 illustrates the results of this experiment. Linoleic acid induces

transcriptional activation of the VCAM-1 promoter by a redox-sensitive **NF**-**kB** like factor. These results are similar to those observed by the activation of VCAM-1 promotor by cytokines such as TNF-.alpha...

DETDESC:

DETD(60)

Confluent . . . native acrylamide gels, and exposed to autoradiography film at -70.degree. C. for 18 hours. Two bands A and C, representing **NF**-**kB** like binding activity are designated. A weak band B was observed in control (untreated) cells.

US PAT NO: 5,807,884 [IMAGE AVAILABLE] DATE ISSUED: Sep. 15, 1998

L15: 24 of 72

Treatment for atherosclerosis and other cardiovascular and TITLE:

inflammatory diseases INVENTOR: Russell M. Medford, Atlanta, GA

R. Wayne Alexander, Atlanta, GA Sampath Parthasarathy, Atlanta, GA Bobby V. Khan, Dunwoody, GA

Emory University, Atlanta, GA (U.S. corp.) 08/317,399 ASSIGNEE:

APPL-NO:

DATE FILED: Oct. 4, 1994

ART-UNIT: 129

PRIM-EXMR: Peter O'Sullivan

LEGAL-REP: Sherry M. Knowles, JacquelineKing & Spalding Haley

US PAT NO: 5,807,884 [IMAGE AVAILABLE] L15: 24 of 72 US-CL-CURRENT: **514/423**, **489**, **506**, **513**, **517**, **518**, **553**, **561**, **824**, **825**, **826**, **861**, **863**; 530/331; 548/431; 549/16

SUMMARY:

BSUM(3)

. of the regulatory elements on the human VCAM-1 gene Molecular . . that control its expression suggests an important role for nuclear factor-kB (**NF**-**kB**), a transcriptional regulatory factor, or an NF-k beta. like binding protein in oxidation-reduction-sensitive regulation of VCAM-1 gene expression. Transcriptional factors are. role in mediating inflammatory and other stress signals to the nuclear regulatory apparatus. Although the precise biochemical signals that activate **NF**-**kB** are unknown, this transcriptional factor may integrate into a common molecular pathway many of the risk factors and "causative" signals. . .

SUMMARY:

BSUM(4)

Importantly, the activation of **NF**-**kB** in vascular endothelial cells by diverse signals can be specifically inhibited by antioxidants such as N-acetylcysteine and pyrrolidine dithiocarbamate (see. . . 07/969,934, now allowed). This has led to the hypothesis that oxygen radicals play an important role in the activation of **NF**-**kB through an undefined oxidation-reduction mechanism. Because an **NF**-**kB**-like enhancer element also regulates the transcription of the VCAM-1 promoter in an oxidation-reduction-sensitive manner, oxidative stress in the atherosclerotic lesion. . .

DRAWING DESC:

DRWD(8)

an illustration of an autoradiogram that indicates that linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive **NF**.***kB** like factor. HAEC were split at the ratio to give approximately 60% confluence in 100-mm tissue culture plates. HAEC were. . .

DRAWING DESC:

DRWD(9)

FIG. 8 is an illustration of an acrylamide gel slab that indicates that polyunsaturated fatty acids activate **NF**-**kB**-like DNA binding polyunsaturated ratty acids activate ""NF"-""*". "RE"-"Ike DNA binding activities that are blocked by the antioxidant PDTC. Confluent HAEC in media containing 4% FBS (as described in. . . native acrylamide gels, and exposed to autoradiography film at -70.degree. C. for 18 hours. Two bands A and C, representing ""NF""-"" [like binding activity are designated. A weak band B was observed in control (untreated) cells.

DETDESC:

DETD(31)

promoter studies that cytokines and non-cytokines activate VCAM-I gene expression in endothelial cells at least in part transcriptionally through two **NF**-**kB**-like DNA binding elements. It has also been demonstrated that PDTC inhibits VCAM-I gene expression through a redox-sensitive **NF**-**kB** like factor. To determine whether polyunsaturated fatty acids induce transcriptional activation of the human VCAM-1 promoter via a similar mechanism,. . . results v obtained with the minimal cytokine-inducible promoter of the VCAM-1 gene (p85 VCAM-CAT), containing the -77 and -63 bp **NF**-**kB**-like sites. Neither linoleic acid nor TNF-.alpha. had any effect on activity using a constitutively expressed pSV.sub.2 CAT construct. PDTC inhibited. indicate that analogous to TNF-.alpha., polyunsaturated fatty acids such as linoleic acid induce the transcriptional activation of VCAM-1 through an **NF**-**kB**-like redox-sensitive mechanism.

DETDESC:

DETD(32)

To determine whether polyunsaturated fatty acids and their oxidative metabolites regulate VCAM-1 promoter activity through an **NF**-**kB**-like transcriptional regulatory factor, nuclear extracts from HAEC were assayed for DNA binding activity to a double-stranded oligonucleotide containing the VCAM-1 **NF**-**kB**-like promoter elements located at positions -77 and -63. As shown in FIG. 7, two bands A and C,*representing **NF**-**kB**-like activity were induced in response to a three hour exposure to linoleic acid (7.5 .mu.M). Similar findings were observed on. . exposure to the cytokine TNF-.alpha. (100 U/ml). A weak band B was observed in control (untreated) cells. No induction of **NF**-**kB**-like binding was observed with the monounsaturated fatty acid oleic acid. Pretreatment of the cells for that PDTC blocks the activation of VCAM-1 gene expression in HUVEC by inhibiting the activation of these **NF** **KB**-like DNA binding proteins.

DETDESC:

DETD(55)

Linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive **NF**-**kB** like factor.

DETDESC:

DETD(57)

FIG. 7 illustrates the results of this experiment. Linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive
NF-**kB** like factor. These results are similar to those observed by the activation of VCAM-1 promotor by cytokines such as TNF-.alpha...

DETDESC:

DETD(59)

Polyunsaturated Fatty Acids Activate **NF**-**kB**-like DNA Binding Activities that are Blocked by the Antioxidant PDTC.

DETDESC:

DETD(60)

Confluent . . . native acrylamide gels, and exposed to autoradiography film at -70.degree. C. for 18 hours. Two bands A and C, representing **NF**-**kB** like binding activity are designated. A weak band B was observed in control (untreated) cells.

DETDESC:

DETD(61)

FIG. 8 illustrates that linoleic acid induces **NF**-**kB** binding activity to VCAM-1 promotor in a redox-sensitive manner. This is analogous to cytokine TNF-.alpha. and suggests a similar mechanism.

US PAT NO: 5,807,746 [IMAGE AVAILABLE]

L15: 25 of 72

DATE ISSUED: Sep. 15, 1998 Method for importing biologically active molecules into TITLE:

cells INVENTOR: Yao-Zhong Lin, Nashville, TN

Jack J. Hawiger, Nashville, TN E: Vanderbilt University, Nashville, TN (U.S. corp.) ASSIGNEE: 08/258,852 APPL-NO:

DATE FILED: Jun. 13, 1994 ART-UNIT: 185

PRIM-EXMR: Nancy Degen

Needle & Rosenberg LEGAL-REP:

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measured by. . .
US PAT NO: 5,801,195 [IMAGE AVAILABLE]
DATE ISSUED: Sep. 1, 1998
                                                                         L15: 26 of 72
DATE ISSUED: Sep. 1, 1998
ITITLE: Immunotherapeutic aryl amides
INVENTOR: George W. Muller, Bridgewater, NJ
Mary Shire, North Plainfield, NJ
David I. Stirling, Branchburg, NJ
                  Celgene Corporation, Warren, NJ (U.S. corp.)
08/366,618
ASSIGNEE:
APPL-NO:
DATE FILED: Dec. 30, 1994
PRIM-EXMR: In-
                   Jane Fan
                    Mathews, Collins, Shepherd & Gould, P.A.
US PAT NO: 5,801,195 [IMAGE AVAILABLE] L15: 26 of 72 US-CL-CURRENT: **514/539**, **532**, **534**, **616**, **617**, **619**, **622**; 560/39, 41, 42; **564/155**, **158**, **169**, **170**, **176**, **180**, **182**, **183**, **219**, **220**
SUMMARY:
BSUM(14)
 The **nuclear** **factor** . **kappa** . **B** (NF. **kappa** . B) is a
pleiotropic transcriptional activator (Lenardo, et al. Cell 1989, 58,
227-29). NF.kappa.B has been implicated as a transcriptional activator.
US PAT NO: 5,798,368 [IMAGE AVAILABLE]
DATE ISSUED: Aug. 25, 1998
TITLE: Tetrasubstituted 2-(2,6-dioxopiperidin-3-yl)-1-
                                                                          L15: 27 of 72
               oxoisoindolines and method of reducing TNF.alpha. levels
            OR: George W. Muller, Bridgewater, NJ
David I. Stirling, Branchburg, NJ
Roger Shen-Chu Chen, Edison, NJ
INVENTOR:
ASSIGNEE:
                   Celgene Corporation, Warren, NJ (U.S. corp.)
APPL-NO: 08/701,494
DATE FILED: Aug. 22, 1996
ART-UNIT: 164
PRIM-EXMR: Jam
                   James H. Reamer
                    Mathews, Collins, Shepherd & Gould, P.A.
LEGAL-REP:
US PAT NO: 5,798,368 [IMAGE AVAILABLE]
                                                                          L15: 27 of 72
US-CL-CURRENT: **514/323**; 546/201
SUMMARY:
BSUM(11)
 The **nuclear** **factor** .**kappa**.**B** (NF.**kappa**.B) is a
pleiotropic transcriptional activator (Lenardo, et al. Cell 1989, 58,
227-29). NF.kappa.B has been implicated as a transcriptional activator.
US PAT NO: 5,795,876 [IMAGE AVAILABLE]
                                                                         L15: 28 of 72
DATE ISSUED: Aug. 18, 1998
TITLE: Method of inhibiting vascular cell adhesion molecule-1 and
               treating chronic inflammatory diseases with 2,
            6-di-alkyl-4-silyl-phenols

OR: Paul S. Wright, Cincinnati, OH

Steven J. Busch, West Chester, OH
INVENTOR:
ASSIGNEE: Hoechst Marion Rousssel, Inc., Cincinnati, OH (U.S. corp.)
 APPL-NO:
                  08/824,221
DATE FILED: Mar. 25, 1997
ART-UNIT: 164
PRIM-EXMR: Phyllis G. Spivack
                    William R. Boudreaux, David M. Stemerick
US PAT NO: 5,795,876 [IMAGE AVAILABLE] US-CL-CURRENT: **514/63**
                                                                          L15: 28 of 72
```

US PAT NO: 5,807,746 [IMAGE AVAILABLE] L15: 25 of US-CL-CURRENT: 435/375; **514/1**, **2**, **21**; 530/300, 350

Having . . . attached to the amino-terminal hydrophobic sequence conferring membrane-permeable capacity. For this purpose a sequence representing a functional domain of the **nuclear** **factor**

"*kappa * . * "B*" (NF-. * kappa * * B) responsible for a nuclear localization signal was selected. Import of such a peptide into the cell would be

DETDESC:

DETD(70)

SUMMARY:

1.15:25 of 72

RSUM(6)

The . . . have been cloned and characterized. For example, both promoters contain multiple DNA sequence elements which can bind the transcription factor, **NF**-**kB**. lademarco, M. F. et al., J. Biol. transcription factor, "NP"---"KB"-, fademarco, M. F. et al., J. Biol. Chem. 267, 16323-16329 (1992); Voraberger, G. et al., J. Immunol. 147, 2777-2786 (1991). The "NP"-"*kB"* family of transcription factors is central in the regulation of several genes upregulated within sites of inflammation. The activation of "NF"-"*kB"* as a transcription factor involves dissociation from an inhibitory subunit, IkB, in the cytoplasm. **NF**-**kB** subunits translocate to the nucleus, bind to specific DNA sequence elements, and activate transcription of several genes, including

```
US PAT NO: 5,792,787 [IMAGE AVAILABLE]
DATE ISSUED: Aug. 11, 1998
              Treatment for atherosclerosis and other cardiovascular and inflammatory diseases
TITLE:
INVENTOR: Russell M. Medford, Atlanta
R. Wayne Alexander, Atlanta, GA
                  Russell M. Medford, Atlanta, GA
            Margaret K. Offermann, Atlanta, GA
ASSIGNEE:
                 Emory University, Atlanta, GA (U.S. corp.)
08/486,239
APPL-NO:
DATE FILED: Jun. 7, 1995
ART-UNIT: 129
PRIM-EXMR:
                   Peter O'Sullivan
LEGAL-REP:
                  Sherry M. Knowles, JacquelineKing & Spalding Haley
US PAT NO: 5.792,787 [IMAGE AVAILABLE]
                                                                       L15: 29 of 72
US-CL-CURRENT: "$14423**, **210**, **315**, **478**, **479**, **484**, **485**, **487**, **488**, **489**, **506**, **513**, **824**, **825**, **826**, **861**, **863**; 546/245;
             548/531, 953; 558/230, 235
DETDESC:
```

DETD(28)

At the molecular level, PDTC has been shown to inhibit the activation of the transcriptional regulatory factor **Nf**-**kB** in response to the transcriptional regulatory factor with the state of the response to certain cytokine and non-cytokine stimuli (Schreck, Rieber et al., 1991; Schreck, Meier et al., 1992). However, by. . . has been discovered that endothelial cells activate VCAM-I gene expression through an apparently novel transcriptional regulatory factor that is not **Nf**-**kB**. This suggests that PDTC may regulate endothelial cell gene expression through its effect on a new transcriptional regulatory protein. It. . .

```
US PAT NO: 5,783,596 [IMAGE AVAILABLE]
DATE ISSUED: Jul. 21, 1998
                                                     L15: 30 of 72
TITLE:
            Treatment for atherosclerosis and other cardiovascular and
inflammatory diseases
INVENTOR: Russell M. Medford, Atlanta, GA
         Margaret K. Offermann, Atlanta, GA
         R. Wayne Alexander, Atlanta, GA
Sampath Parthasarathy, Atlanta, GA
              Emory University, Atlanta, GA (U.S. corp.)
ASSIGNEE:
APPL-NO:
             08/477,881
DATE FILED: Jun. 7, 1995
ART-UNIT:
              129
PRIM-EXMR:
              Peter O'Sullivan
LEGAL-REP:
              Sherry M. Knowles, JacquelineKing & Spalding Haley
US PAT NO: 5 783 596 HMAGE AVAILABLEL
                                                     1.15: 30 of 72
```

DETDESC:

DETD(56)

At the molecular level, PDTC has been shown to inhibit the activation of the transcriptional regulatory factor **Nf**-**kB** in response to the transcriptional regulatory factor "Net"—Seb" in response to certain cytokine and non-cytokine stimuli (Schreck, Rieber et al. 1991; Schreck, Meier et al. 1992). However, by. . . has been discovered that endothelial cells activate VCAM-1 gene expression through an apparently novel transcriptional regulatory factor that is not **Nf**-**kB**. This suggests that PDTC may regulate endothelial cell gene expression through its effect on a new transcriptional regulatory protein. It.

US PAT NO: 5,783,565 [IMAGE AVAILABLE] DATE ISSUED: Jul. 21, 1998 L15: 31 of 72

TITLE: Cationic amphiphiles containing spermine or spermidine

cationic group for intracellular delivery of therapeutic

molecules

INVENTOR: Edward R. Lee, Quincy, MA

David J. Harris, Lexington, MA Craig S. Siegel, Woburn, MA Seng H. Cheng, Wellesley, MA

Simon J. Eastman, Marlboro, MA John Marshall, Milford, MA Ronald K. Scheule, Hopkinton, MA

ASSIGNEE: Genzyme Corporation, Framingham, MA (U.S. corp.)

APPL-NO: 08/595.375 DATE FILED: Feb. 1, 1996

ART-UNIT:

PRIM-EXMR: Jasemine C. Chambers ASST-EXMR:

Abdur Razzaque Finnegan, Henderson, Farabow, Garrett & Dunner, L.L.P. LEGAL-REP:

US PAT NO: 5,783,565 [IMAGE AVAILABLE] US-CL-CURRENT: **514/44**; 424/450; 536/23.1; 552/544 L15: 31 of 72

DETDESC:

DETD(297)

It . . . increase with the severity of an inflammatory condition (for example, tumor necrosis factor "TNF" and potentially transcription factors such as **NF**-**kB**, AP-1, NF-IL6 and octamer binding protein). It has also been determined that interleukin 8, a polypeptide of 8,500 MW, is. . .

US PAT NO: 5,780,220 [IMAGE AVAILABLE]

L15: 32 of 72

TITLE: Methods and compositions for inhibiting HIV replication INVENTOR: David B. Weiner Merion D4

Yosef Refaeli, Boston, MA

David N. Levy, Birmingham, AL

Trustees of the University of Pennsylvania, Philadelphia, ASSIGNEE:

PA (U.S. corp.) APPL-NO: 08/382,873

DATE FILED: Feb. 3, 1995 ART-UNIT: 188

PRIM-EXMR: ASST-EXMR:

Jeffrey S. Parkin LEGAL-REP:

Woodcock Washburn Kurtz Mackiewicz & Norris, LLP

US PAT NO: 5,780,220 [IMAGE AVAILABLE] L15: 32 of 72 US-CL-CURRENT: 435/5; 424/188.1; 435/7.1; **514/49**, **51**, **179**; 530/350

SUMMARY:

BSUM(10)

The . . . infection of myeloid cell lines can result in a more differentiated phenotype and increase the expression of factors such as **NF**.**KB** which are necessary for HIV replication. Roulston, A. et al. (1992) J. Exp. Med. 175:751; and Chantal Petit, A. J. . .

US PAT NO: 5,773,231 [IMAGE AVAILABLE]

L15: 33 of 72

DATE ISSUED: Jun. 30, 1998

Treatment for atherosclerosis and other cardiovascular and inflammatory diseases R: Russell M. Medford, Atlanta, GA

INVENTOR:

R. Wayne Alexander, Atlanta, GA Sampath Parthasarathy, Atlanta, GA

Bobby V. Khan, Dunwoody, GA ASSIGNEE: Emory University, Atlanta, GA (U.S. corp.)

APPL-NO: 08/473,272 DATE FILED: Jun. 7, 1995

129 ART-UNIT: PRIM-EXMR: Peter O'Sullivan

Sherry M.King & Spalding Knowles LEGAL-REP:

235; **564/76**; 568/21, 25

5,773,231 [IMAGE AVAILABLE] L15: 33 of 72 US PAT NO: US-CL-CURRENT: 4357.24; "*14/489", "*506", "*513", "*825"*, **861", "*863"; 530/331; 548/431; 558/230,

SUMMARY:

BSUM(4)

Molecular . . . of the regulatory elements on the human VCAM-1 gene that control its expression suggests an important role for nuclear factor-kB (**NF**-**kB**), a transcriptional regulatory factor, or an NF-k.beta. like binding protein in oxidation-reduction-sensitive regulation of VCAM-1 gene expression. Transcriptional factors are. . .

role in mediating inflammatory and other stress signals to the nuclear regulatory apparatus. Although the precise biochemical signals that activate **NF**-**kB** are unknown, this transcriptional factor may integrate into a common molecular pathway many of the risk factors and "causative" signals. . .

SUMMARY:

BSUM(5)

Importantly, the activation of **NF**-**kB** in vascular endothelial cells by diverse signals can be specifically inhibited by antioxidants such as N-acetylcysteine and pyrrolidine dithiocarbamate (see. . 07/969,934, now allowed). This has led to the hypothesis that oxygen radicals play an important role in the activation of **NF**-**kB** through an undefined oxidation-reduction mechanism. Because an **NF**-**kB**-like enhancer element also regulates the transcription of the VCAM-1 promoter in an oxidation-reduction-sensitive manner, oxidative stress in the atherosclerotic lesion. . .

DRAWING DESC:

DRWD(8)

FIG. . . . an illustration of an autoradiogram that indicates that linoleic acid induces transcriptional activation of the VCAM-I promoter by a redox-sensitive **NF**-**kB** like factor. HAEC were split at the ratio to give approximately 60% confluence in 100-mm tissue culture plates. HAEC were. . .

DRAWING DESC:

DRWD(9)

FIG. 8 is an illustration of an acrylamide gel slab that indicates that polyunsaturated fatty acids activate **NF**-**kB**-like DNA binding activities that are blocked by the antioxidant PDTC. Confluent HAEC in activities that are proceed by the annovation in Parts. Communit Parts in media containing 4% FBS (as described in. . . native acrylamide gels, and exposed to autoradiography film at -70.degree. C. for 18 hours. Two bands A and C, representing **NF**.**kB** like binding activity are designated. A weak band B was observed in control (untreated) cells.

DETDESC:

DETD(31)

Previous . . . promoter studies that cytokines and non-cytokines activate VCAM-1 gene expression in endothelial cells at least in part transcriptionally through two **NF**-**kB**-like DNA binding elements. It has also been demonstrated that PDTC inhibits VCAM-1 gene expression through a redox-sensitive **NF**_**kB** like factor. To determine whether polyunsaturated fatty acids induce transcriptional activation of the human VCAM-I promoter via a similar mechanism, . . . results were obtained with the minimal cytokine-inducible promoter of the VCAM-I gene (p85 VCAM-CAT), containing the -77 and -63 bp **NF**-**kB**-like sites. Neither linoleic acid nor TNF-.alpha. had any effect on activity using a constitutively expressed pSV.sub.2 CAT construct. PDTC inhibited. indicate that analogous to TNF-.alpha., polyunsaturated fatty acids such as linoleic acid induce the transcriptional activation of VCAM-1 through an **NF**-**kB**-like redox-sensitive mechanism.

DETDESC:

DFTD(32)

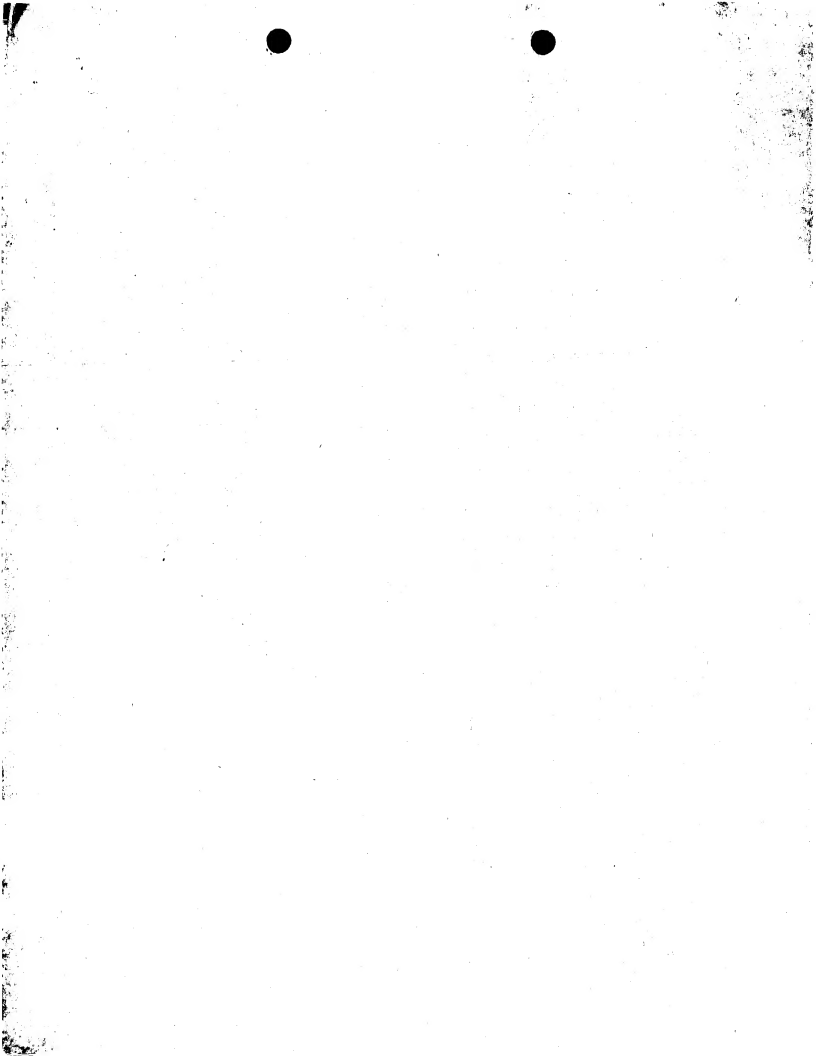
To determine whether polyunsaturated fatty acids and their oxidative metabolites regulate VCAM-1 promoter activity through an
NF-**kB**-like transcriptional regulatory factor, nuclear extracts
from HAEC were assayed for DNA binding activity to a double-stranded oligonucleotide containing the VCAM-1 **NF**-**kB**-like promoter oligonucleotide containing the VCAM-1 **NF***-- **NB**---- inc promoter between the located at positions -77 and -63. As shown in FIG. 7, two bands A and C, representing **NF**-******--like activity were induced in response to a three hour exposure to linoleic acid (7.5 .mu. M). Similar findings were observed on. . . exposure to the cytokine TNF-.alpha (100 U/ml). A weak band B was observed in control (untreated) cells. No induction of **NF**-***kB**-like binding was observed with the monounsaturated fatty acid oleic acid. Pretreatment of the cells for thirty minutes with PDTC inhibited. . . previously reported findings that PDTC blocks the activation of VCAM-1 gene expression in HUVEC by inhibiting the activation of these **NF** **KB**-like DNA binding proteins.

DETDESC:

DETD(55)

Linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive **NF**-**kB** like factor.

DETDESC:



DETD(57)

FIG. 7 illustrates the results of this experiment. Linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive **NF**-**kB** like factor. These results are similar to those observed by the activation of VCAM-1 promotor by cytokines such as TNF-.alpha...

DETDESC:

DETD(59)

Polyunsaturated Fatty Acids Activate **NF**-**kB**-like DNA Binding Activities that are Blocked by the Antioxidant PDTC.

DETDESC:

DETD(60)

Confluent . . . native acrylamide gels, and exposed to autoradiography film at -70.degree. C. for 18 hours. Two bands A and C, representing **NF**.**kB** like binding activity are designated. A weak band B was observed in control (untreated) cells.

DETDESC:

DETD(61)

FIG. 8 illustrates that linoleic acid induces **NF**-**kB** binding activity to VCAM-1 promotor in a redox-sensitive manner. This is analogous to cytokine TNF-.alpha. and suggests a similar mechanism.

US PAT NO: 5,773,209 [IMAGE AVAILABLE] DATE ISSUED: Jun. 30, 1998

L15: 34 of 72

Treatment for atherosclerosis and other cardiovascular and

inflammatory diseases
R: Russell M. Medford, Atlanta, GA INVENTOR: R. Wayne Alexander, Atlanta, GA

Sampath Parthasarathy, Atlanta, GA Bobby V. Khan, Dunwoody, GA

ASSIGNEE: Emory University, Atlanta, GA (U.S. corp.)

APPL-NO: 08/484,059 DATE FILED: Jun. 7, 1995 ART-UNIT: PRIM-EXMR: 129 Peter O'Sullivan

LEGAL-REP:

Sherry M.King & Spalding Knowles

5.773.209 (IMAGE AVAILABLE) L15: 34 of 72 US-CL-CURRENT: 435/7.24; 424/9.1, 9.2; 435/6, 7.2, 7.21, 7.94, 7.95; 436/71, 86, 129, 172, 503, 504, 548; **514/18**, **423**, **478**, **478**, **484**, **485**, **487**,

SUMMARY:

BSUM(5)

Molecular . . . of the regulatory elements on the human VCAM-1 gene that control its expression suggests an important role for nuclear factor-kB (**NF**-**kB**), a transcriptional regulatory factor, or an NF-k.beta. like binding protein in oxidation-reduction-sensitive regulation of VCAM-1 gene expression. Transcriptional factors are. . role in mediating inflammatory and other stress signals to the nuclear regulatory apparatus. Although the precise biochemical signals that *NF**-**kB** are unknown, this transcriptional factor may integrate into a common molecular pathway many of the risk factors and "causative" signals.

SUMMARY:

BSUM(6)

Importantly, the activation of **NF**-**kB** in vascular endothelial cells by diverse signals can be specifically inhibited by antioxidants such as N-acetylcysteine and pyrrolidine dithiocarbamate (see. through an undefined oxidation-reduction mechanism. Because an **NF**-**kB**-like enhancer element also regulates the transcription of the VCAM-1 promoter in an oxidation-reduction-sensitive manner, oxidative stress in the atherosclerotic lesion. . .

DRAWING DESC:

FIG. . . . an illustration of an autoradiogram that indicates that linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive **NF**-**kB** like factor. HAEC were split at the ratio to give approximately 60% confluence in 100-mm tissue culture plates. HAEC were.

DRAWING DESC:

DRWD(9)

FIG. 8 is an illustration of an acrylamide gel slab that indicates that polyunsaturated fatty acids activate **NF**-**kB**-like DNA binding activities that are blocked by the antioxidant PDTC. Confluent HAEC in designated. A weak band B was observed in control (untreated) cells.

DETDESC:

DETD(31)

. promoter studies that cytokines and non-cytokines activate VCAM-I gene expression in endothelial cells at least in part transcriptionally through two **NF**-**kB**-like DNA binding elements. It has also been demonstrated that PDTC inhibits VCAM-1 gene expression through a redox-sensitive **NF**-**kB** like factor. To determine whether polyunsaturated fatty acids induce transcriptional activation of the human VCAM-1 promoter via a similar mechanism, results were obtained with the minimal cytokine-inducible promoter of the VCAM-1 gene (p85 VCAM-CAT), containing the -77 and -63 bp **NF**-**kB**-like sites. Neither linoleic acid nor TNF. alpha, had any effect on activity using a constitutively expressed pSV.sub.2 CAT construct. PDTC inhibited. indicate that analogous to TNF-.alpha., polyunsaturated fatty acids such as linoleic acid induce the transcriptional activation of VCAM-1 through an **NF**-**kB**-like redox-sensitive mechanism.

DETDESC:

DETD(32)

To determine whether polyunsaturated fatty acids and their oxidative metabolites regulate VCAM-1 promoter activity through an
NF-**kB**-like transcriptional regulatory factor, nuclear extracts from HAEC were assayed for DNA binding activity to a double-stranded oligonucleotide containing the VCAM-1 **NF*--*kB**-like promoter elements located at positions -77 and -63. As shown in FIG. 7, two bands A and C, representing **NF**-**kB**-like activity were induced in response to a three hour exposure to linoleic acid (7.5 .mu.M). Similar findings were observed on. . . exposure to the cytokine TNF-alpha. (100 U/ml). A weak band B was observed in control (untreated) cells. No induction of **NF**.**kB**-like binding was observed with the monounsaturated fatty acid oleic acid. Pretreatment of the cells for that PDTC blocks the activation of VCAM-1 gene expression in HUVEC by inhibiting the activation of these **NF** **KB**-like DNA binding proteins.

DETDESC-

Linoleic Acid Induces Transcriptional Activation of the VCAM-1 Promoter by a Redox-sensitive **NF**-**kB** Like Factor

DETDESC:

DETD(57)

FIG. 7 illustrates the results of this experiment. Linoleic acid induces transcriptional activation of the VCAM-I promoter by a redox-sensitive
NF-**kB** like factor. These results are similar to those observed by the activation of VCAM-1 promotor by cytokines such as TNF-.alpha...

DETDESC:

DETD(59)

Polyunsaturated Fatty Acids Activate **NF**-**kB**-like DNA Binding Activities that are Blocked by the Antioxidant PDTC

DETDESC:

DETD(60)

Confluent . . . native acrylamide gels, and exposed to autoradiography film at -70.degree. C. for 18 hours. Two bands A and C, representing **NF**-**kB** like binding activity are designated. A weak band B was observed in control (untreated) cells.

US PAT NO: 5,770,581 [IMAGE AVAILABLE] DATE ISSUED: Jun. 23, 1998

L15: 35 of 72

TITLE: Gene transcription and ionizing radiation: methods and

compositions

INVENTOR: Ralph R. Weichselbaum, Chicago, IL Dennis E. Hallahan, Park Ridge, IL

Vikas P. Sukhatme, Chicago, IL Donald W. Kufe, Wellesley, MA

ASSIGNEE: Arch Development Corp., Chicago, IL (U.S. corp.) Dana-Farber Cancer Institute, Boston, MA (U.S. corp.)

APPL-NO: 08/474,445

DATE FILED: Jun. 7, 1995

ART-UNIT:

PRIM-EXMR: Bruce R. Campbell LEGAL-REP: Arnold, White & Durkee

US PAT NO: 5,770,581 [IMAGE AVAILABLE] L15: 35 of 72

US-CL-CURRENT: **514/44**; 435/447, 455; 536/24.1

DETDESC:

DETD(85)

Transcription . . . domains are well known in the art. Exemplary transcription factors having activation domains are GAL4, c-Jun, viral protein VP-16, and **nuclear** **factor** NF-.**kappa**.**B**.

DETDESC:

DETD(100)

Nuclear **factor** NF-.**kappa**.**B** is a transcription factor. The activation domain of NF-.kappa.B comprises amino acid residue sequences from about residue position 414 to. . .

DETDESC:

DETD(272)

The . . . al., 1990; Hallahan, et al, 1991). Other studies have demonstrated that x-rays induce expression and DNA binding activity of the **nuclear** **factor** .**kappa**.**B** (NF-.**kappa**.B; Brach, et

DETDESC:

DETD(315)

lonizing . . . which code for transcription factors. Other studies have demonstrated that ionizing radiation induces expression and DNA binding activity of the **nuclear** **factor** .**kappa**.**B** (NF-.**kappa**.B). The activation of transcription factors likely represents a critical control point in transducing early nuclear signals to longer term changes. . .

DETDESC:

DETD(330)

NAC . . . phorbol ester-induced activation of the HIV-1 long terminal repeat. This antioxidant has also been found to inhibit activation of the **nuclear** **factor** .**kappa**.**B** (NF-.**kappa**.B) by phorbol esters and other agents such as H.sub.2 O.sub.2. The available findings suggest the release activate NF-.kappa.B by induced. . .

US PAT NO: 5,767,099 [IMAGE AVAILABLE] DATE ISSUED: Jun. 16, 1998

L15: 36 of 72

Cationic amphiphiles containing amino acid or dervatized TITLE:

amino acid groups for intracellular delivery of

therapeutic molecules

INVENTOR: David J. Harris, Lexington, MA

JR: David J. Harris, Lexington, Edward R. Lee, Quincy, MA Craig S. Siegel, Woburn, MA Eric A. Rowe, Malden, MA Shirley C. Hubbard, Belmont, MA

Genzyme Corporation, Cambridge, MA (U.S. corp.) ASSIGNEE:

APPL-NO: 08/546,086

DATE FILED: Oct. 20, 1995

184 ART-UNIT:

PRIM-EXMR: Jacqueline M. Stone Patrick Twomey ASST-EXMR:

LEGAL-REP: E. Victor Donahue

US PAT NO: 5,767,099 [IMAGE AVAILABLE] L15: 36 of 72 US-CL-CURRENT: **514/44**, **182**, **777**; 516/915; 552/544; 560/6

DETDESC:

DETD(297)

It . . . increase with the severity of an inflammatory condition (for

example, tumor necrosis factor "TNF" and potentially transcription factors such as ""NF"-""kB", AP-1, NF-IL6 and octamer binding protein). It has also been determined that interleukin 8, a polypeptide of 8,500

US PAT NO: 5,750,351 [IMAGE AVAILABLE]

L15: 37 of 72

DATE ISSUED: May 12, 1998

Treatment for atherosclerosis and other cardiovascular and TITLE:

inflammatory diseases INVENTOR:

Russell M. Medford, Atlanta, GA R. Wayne Alexander, Atlanta, GA Sampath Parthasarathy, Atlanta, GA Bobby V. Khan, Dunwoody, GA

ASSIGNEE: Emory University, Atlanta, GA (U.S. corp.)

08/474,530 APPL-NO:

DATE FILED: Jun. 7, 1995

ART-UNIT: 129

Peter O'Sullivan PRIM-EXMR:

LEGAL-REP: Sherry M. Knowles, JacquelineKing & Spalding Haley

US PAT NO: 5,750,351 [IMAGE AVAILABLE] 1.15: 37 of 72 US PAT NO: 5,750,351 [IMAGE AVAILABLE] LIS: 37 07 12
US-CL-CURRENT: 435/7.21; 424/9.1, 9.2; 435/6, 7.2, 7.24, 7.94, 7.95;
436/71, 86, 129, 172, 503, 504, 548; "514/18"*,
"226,2"*, "423"*, "477"*, "478"*, "479"*, "484"*,
"485"*, "487"*, "488"*, "489"*, "506"*, "513"*,
"517"*, "518"*, "553"*, "561"*, "824"*, "825"*,
"826"*, "861"*, "863"*; 530/331; 548/431; 549/16;
558/230, 234, 235, 250; 562/26, 27; "564/76"*

SUMMARY:

BSUM(4)

Molecular . . . of the regulatory elements on the human VCAM-I gene factor-kB (**NF**-**kB**), a transcriptional regulatory factor, or an NF-k.beta. like binding protein in oxidation-reduction-sensitive regulation of VCAM-1 gene expression. Transcriptional factors are. . role in mediating inflammatory and other stress signals to the nuclear regulatory apparatus. Although the precise biochemical signals that activate **NF**-**kB** are unknown, this transcriptional factor may integrate into a common molecular pathway many of the risk factors and "causative" signals. . .

SUMMARY: BSUM(5)

Importantly, the activation of **NF**-**kB** in vascular endothelial cells by diverse signals can be specifically inhibited by antioxidants such as N-acetylcysteine and pyrrolidine dithiocarbamate (see. . . 07/969,934, now allowed). This has led to the hypothesis that oxygen radicals play an important role in the activation of **NF**-**kB** **NF**-**kB**-like enhancer element also regulates the transcription of the VCAM-1 promoter in an oxidation-reduction-sensitive manner, oxidative stress in the atherosclerotic lesion. . .

DRAWING DESC:

DRWD(8)

an illustration of an autoradiogram that indicates that linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive **NF**.**kB** like factor. HAEC were split at the ratio to give approximately 60% confluence in 100-mm tissue culture plates. HAEC were. .

DRAWING DESC:

DRWD(9)

FIG. 8 is an illustration of an acrylamide gel slab that indicates that polyunsaturated fatty acids activate **NF**-**kB**-like DNA binding polyuisaturiate ratty activities that are blocked by the antioxidant PDTC. Confluent HAEC in media containing 4% FBS (as described in. . . native acrylamide gels, and exposed to autoradiography film at -70.degree. C. for 18 hours. Two bands A and C, representing **NF**-*kB** like binding activity are designated. A weak band B was observed in control (untreated) cells.

DETDESC:

DETD(31)

. promoter studies that cytokines and non-cytokines activate VCAM-1 gene expression in endothelial cells at least in part transcriptionally through two **NF**-**kB**-like DNA binding elements. It has also been demonstrated that PDTC inhibits VCAM-1 gene expression through a redox-sensitive **NF**-**kB** like factor. To determine whether

polyunsaturated fatty acids induce transcriptional activation of the human VCAM-1 promoter via a similar mechanism... results were obtained with the minimal cytokine-inducible promoter of the VCAM-1 gene (p85 VCAM-CAT), containing the -77 and -63 bp **NF**=*kB**-like sites. Neither linoleic acid nor TNF- alpha. had any effect on activity using a constitutively expressed pSV.sub.2 CAT construct. PDTC inhibited. . . indicate that analogous to TNF-alpha., polyunsaturated fatty acids such as linoleic acid induce the transcriptional activation of VCAM-1 through an **NF**-**kB**-like redox-sensitive mechanism.

DETDESC:

DETD(32)

To determine whether polyunsaturated fatty acids and their oxidative metabolites regulate VCAM-1 promoter activity through an **NF**-**kB**-like transcriptional regulatory factor, nuclear extracts from HAEC were assayed for DNA binding activity to a double-stranded oligonucleotide containing the VCAM-1 "*NF*--*kB*-like promoter elements located at positions -77 and -63. As shown in FIG. 7, two bands A and C, representing **NF*--**kB*-like activity were induced in response to a three hour exposure to linoleic acid (7.5 .mu.M). Similar findings were observed on. . . exposure to the cytokine TNF- alpha. (100 U/ml). A weak band B was observed in control (untreated) cells. No induction of **NF**.**kB**-like binding was observed with the monounsaturated fatty acid oleic acid. Pretreatment of the cells for thirty minutes with PDTC inhibited. . . previously reported findings that PDTC blocks the activation of VCAM-1 gene expression in HUVEC by inhibiting the activation of these **NF** **KB**-like DNA binding proteins.

DETDESC:

DETD(57)

FIG. 7 illustrates the results of this experiment. Linoleic acid induces transcriptional activation of the VCAM-1 promoter by a redox-sensitive **NF**-**kB** like factor. These results are similar to those observed by the activation of VCAM-1 promotor by cytokines such as TNF-.alpha...

DETDESC:

DETD(59)

Polyunsaturated Fatty Acids Activate **NF**-**kB**-like DNA Binding Activities that are Blocked by the Antioxidant PDTC

DETDESC:

DETD(60)

Confluent . . . native acrylamide gels, and exposed to autoradiography film at -70.degree. C. for 18 hours. Two bands A and C, representing **NF**-**kB** like binding activity are designated. A weak band B was observed in control (untreated) cells.

DETDESC:

DETD(61)

FIG. 8 illustrates that linoleic acid induces **NF**-**kB** binding activity to VCAM-1 promotor in a redox-sensitive manner. This is analogous to cytokine TNF-.alpha. and suggests a similar mechanism. . .

5,747,471 [IMAGE AVAILABLE] US PAT NO: DATE ISSUED: May 5, 1998

Cationic amphiphiles containing steroid lipophilic groups for intracellular delivery of therapeutic molecules Craig S. Siegel, Woburn, MA TITLE:

David J. Harris, Lexington, MA Edward R. Lee, Quincy, MA Shirley C. Hubbard, Belmont, MA Seng H. Cheng, Wellesley, MA Simon J. Eastman, Marlboro, MA John Marshall, Milford, MA Ronald K. Scheule, Hopkinton, MA

Mathieu B. Lane, Cambridge, MA

Eric A. Rowe, Malden, MA Genzyme Corporation, Cambridge, MA (U.S. corp.) 08/540,867 ASSIGNEE:

APPL-NO: DATE FILED: Oct. 11, 1995 ART-UNIT: 184

Jacqueline M. Stone PRIM-EXMR: ASST-EXMR: Patrick Twomey E. Victor Donahue LEGAL-REP:

US PAT NO: 5,747,471 [IMAGE AVAILABLE] L15: 38 US-CL-CURRENT: **514/44**, **182**, **777**; 552/544; 560/6 L15: 38 of 72

DETDESC:

DETD(296)

It . . . that increase with the severity of an inflammatory condition (for example, tumor necrosis factor "TNF", and transcription factors such as "*NF". ***** AP-1, NF-IL6 and octamer binding protein). It has also been determined that interleukin 8, a polypeptide of 8,500 MW, is.

US PAT NO: 5,744,131 [IMAGE AVAILABLE] DATE ISSUED: Apr. 28, 1998

L15: 39 of 72

TITLE: Sequence-directed DNA-binding molecules compositions and methods

INVENTOR: Cynthia A. Edwards, Menlo Park, CA

Kirk E. Fry, Palo Alto, CA Charles R. Cantor, Boston, MA

Beth M. Andrews, Maynard, MA

Genelabs Technologies, Inc., Redwood City, CA (U.S. corp.) ASSIGNEE:

APPL-NO: 08/476,876 DATE FILED: Jun. 7, 1995 ART-UNIT:

PRIM-EXMR: Stephanie W. Zitomer

ASST-EXMR: Amy Atzel

Gary R. Fabian, Carol A. Stratford, Peter J. Dehlinger LEGAL-REP:

US PAT NO: 5,744,131 [IMAGE AVAILABLE]

L15: 39 of 72

US-CL-CURRENT: 424/78.08; 436/501; **514/1**

DETDESC:

DETD(219)

Similarly, . . . nuclear factor (HNF-1), which is required for the expression of human hepatitis B virus (HBV) (Chang, H. -K.), and (ii) **NFkB** and NFAT-1 binding sites in the human immunodeficiency virus (HIV) long terminal repeat (LTR), one or both of which may. . .

DETDESC:

DETD(222)

of F2 replication

genital warts

cervical carcinoma

Interleukin 2 NFAT-1

immunosuppressant

enhancer HIV LTR NFAT-1 AIDS, ARC

NFkB

HBV enhancer HNF-1 hepatitis

Fibrogen promoter

HNF-1 cardiovascular

disease Oncogene promoter cancer

and coding sequences. . .

DETDESC:

DETD(223)

(Abbreviations: . . virus; HPV, human papilloma virus; HIV LTR, Human immunodeficiency virus long terminal repeat; NFAT, nuclear factor of activated T cells; **NFkB**, nuclear factor kappaB; AIDS, acquired immune deficiency syndrome; ARC, AIDS related complex; HBV, hepatitis
B virus; HNF, hepatic **nuclear** **factor**.)

US PAT NO: 5,738,852 [IMAGE AVAILABLE] DATE ISSUED: Apr. 14, 1998

L15: 40 of 72

Methods of enhancing antigen-specific T cell responses INVENTOR:

William S. Robinson, Palo Alto, CA

Keting Chu, Palo Alto, CA EE: Solis Therapeutics, Inc., Palo Alto, CA (U.S. corp.)

ASSIGNEE: 08/663,157 APPL-NO:

DATE FILED: Jul. 29, 1996

185 ART-UNIT:

Johnny F. Railey, II PRIM-EXMR:

LEGAL-REP: Pennie & Edmonds LLP

US PAT NO: 5,738,852 [IMAGE AVAILABLE] L15: 40 of 72 US-CL-CURRENT: 424/199.1, 93.2, 278.1; 435/320.1; **514/44**

DETDESC:

DETD(37)

In . . . be inserted upstream of the transcriptional control regions.

Alternatively, or in addition, multimeric transcription factor binding sites (e.g., NF-AT and/or **NFKB**) may be inserted into or upstream of the transcriptional control regions, combining the upstream region of one with the proximal. . .

US PAT NO: 5,736,570 [IMAGE AVAILABLE] DATE ISSUED: Apr. 7, 1998

L15: 41 of 72

TITLE: Immunotherapeutic aryl amides
INVENTOR: George W. Muller, Bridgewater, NJ
Mary Shire, North Plainfield, NJ

David I. Stirling, Branchburg, NJ
EE: Celgene Corporation, Warren, NJ (U.S. corp.)
D: 08/729,847 ASSIGNEE:

APPL-NO: DATE FILED: Oct. 15, 1996 ART-UNIT: 123 PRIM-EXMR: Jane Fan

LEGAL-REP: Mathews, Collins, Shepherd & Gould

US PAT NO: 5,736,570 [IMAGE AVAILABLE] L15: 41 of US-CL-CURRENT: **514/532**, **535**, **617**, **619**, **622* L15: 41 of 72

BSUM(14)

The **nuclear** **factor** .**kappa**.**B** (NF. **kappa**.B) is a pleiotropic transcriptional activator (Lenardo, et al. Cell 1989, 58, 227-29). NF.kappa.B has been implicated as a transcriptional activator.

US PAT NO: 5,733,762 [IMAGE AVAILABLE] DATE ISSUED: Mar. 31, 1998

L15: 42 of 72

Complexes of nucleic acid and polymer, their process of preparation and their use for the transfection of cells
R: Patrick Midoux, Orleans, France

INVENTOR:

Patrick Erbacher, Orleans, France

Annie-Claude Roche-Degremont, Sandillon, France

Michel Monsigny, Saint-Cyr-En-Val, France
E: I.D.M. Immuno-Designed Molecules, France (foreign corp.) ASSIGNEE:

08/741,678 DATE FILED: Oct. 31, 1996 ART-UNIT: 189

PRIM-EXMR: George C. Elliott ASST-EXMR: Thomas G. Larson

LEGAL-REP: Bierman, Muserlian and Lucas

US PAT NO: 5,733,762 [IMAGE AVAILABLE] L15: 42 of 72 US-CL-CURRENT: 435/458, 325; **514/44**; 530/300, 345, 350, 395, 402; 536/23.2, 23.5, 23.7, 24.5

SUMMARY:

BSUM(146)

nuclear factors: **NF**-**KB**, CII TA, . . .

US PAT NO: 5,723,335 [IMAGE AVAILABLE]
DATE ISSUED: Mar. 3, 1998 L15: 43 of 72

Immune stimulation by phosphorothioate oligonucleotide

INVENTOR: Stephen L. Hutcherson, Richmond, VA

Josephine M. Glover, Woking, United Kingdom

E: Isis Pharmaceuticals, Inc., Carlsbad, CA (U.S. corp.)

ASSIGNEE:

APPL-NO: 08/712,135 DATE FILED: Sep. 11, 1996 ART-UNIT: 189

PRIM-EXMR: Charles C.P. Rories

LEGAL-REP: Law Offices of Jane Massey Licata

5,723,335 [IMAGE AVAILABLE] L15: 43 of 72 US-CL-CURRENT: 435/375; 424/1.73, 1.77, 280.1; **514/44**; 536/23.1, 24.3, 24.31, 24.33

SUMMARY:

BSUM(18)

Oligonucleotides having a sequence identical to a portion of the sense strand of the mRNA encoding the p65 subunit of **NF**-**kB**, a DNA binding protein, were found to stimulate splenic cell proliferation both in vitro and in vivo. The proliferating spleen cells were shown to be B cells. Immunoglobulin secretion and **NF**-**kB** activity in these cell lines was also increased by the sense oligonucleotide. Both phosphodiester and phosphorothioate sense oligonucleotides stimulated

US PAT NO: 5,719,131 [IMAGE AVAILABLE]

L15: 44 of 72

DATE ISSUED: Feb. 17, 1998

Cationic amphiphiles containing dialkylamine lipophilic groups for intracellular delivery of therapeutic

INVENTOR:

DR: David J. Harris, Lexington, MA Edward R. Lee, Quincy, MA Craig S. Siegel, Woburn, MA Seng H. Cheng, Wellesley, MA Simon J. Eastman, Marlboro, MA John Marshall, Milford, MA

Ronald K. Scheule, Hopkinton, MA

Genzyme Corporation, Framingham, MA (U.S. corp.) 08/546,110 ASSIGNEE:

APPL-NO: DATE FILED: Oct. 20, 1995

ART-UNIT: 184 PRIM-EXMR: Ch

Christopher S.F. Low Dave T. Nguyen ASST-EXMR:

US PAT NO: 5,719,131 [IMAGE AVAILABLE] US-CL-CURRENT: **514/44**; 424/450; 552/544 1.15: 44 of 72

DETDESC:

DETD(297)

It . . . increase with the severity of an inflammatory condition (for example, tumor necrosis factor "TNF" and potentially transcription factors such as **NF**-**kB**, AP-1, NF-IL6 and octamer binding protein). It has also been determined that interleukin 8, a polypeptide of 8,500

US PAT NO: 5,703,098 [IMAGE AVAILABLE] DATE ISSUED: Dec. 30, 1997 L15: 45 of 72

ITTLE: Immunotherapeutic imides/amides
INVENTOR: George W. Muller, Bridgewater, NJ
Mary Shire, North Plainfield, NJ

David I. Stirling, Branchburg, NJ
E: Celgene Corporation, Warren, NJ (U.S. corp.)
: 08/759,788 ASSIGNEE:

APPL-NO: DATE FILED: Dec. 3, 1996 ART-UNIT: 121

PRIM-EXMR: Floyd D. Higel

LEGAL-REP: Mathews, Collins, Shepherd & Gould, P.A.

L15: 45 of 72

US PAT NO: 5,703,098 [IMAGE AVAILABLE] L15 US-CL-CURRENT: **514/339**, **417**; 546/277.1; 548/476

SUMMARY:

BSUM(14)

The **nuclear** **factor** . **kappa** . **B** (NF. **kappa** . B) is a pleiotropic transcriptional activator (Lenardo, et al. Cell 1989, 58, 227-29). NF.kappa.B has been implicated as a transcriptional activator.

US PAT NO: 5,703,069 [IMAGE AVAILABLE] DATE ISSUED: Dec. 30, 1997 L15: 46 of 72

Method for inhibiting and controlling viral growth

INVENTOR: OR: David Thomas Connor, Ann Arbor, MI Stephen Joseph Gracheck, Ann Arbor, MI

ASSIGNEE:

Warner-Lambert Company, Morris Plains, NJ (U.S. corp.)

APPL-NO: 08/712,063 DATE FILED: Sep. 11, 1996

ART-UNIT:

PRIM-EXMR: Robert T. Bond LEGAL-REP: Charles W. Ashbrook

US PAT NO: 5,703,069 [IMAGE AVAILABLE] US-CL-CURRENT: **514/211**, **220**; 540/488, 495 L15: 46 of 72

SUMMARY:

BSUM(7)

An . . . further development is therapeutic targeting along cellular signaling pathways that result in HIV-1 transcriptional activation. Among the potential targets is **nuclear** **factor**-.**kappa** **B** (NF-.**kappa**.B), a transcriptional enhancer important for HIV-1 activation. In resting cells, preformed NF-, kappa. B exists in the cytoplasm bound to its inhibitor.

L15: 47 of 72

US PAT NO: 5,698,579 [IMAGE AVAILABLE] DATE ISSUED: Dec. 16, 1997

Dec. 16, ITTLE: Cyclic amides INVENTOR: George ***

INVENTOR: George W. Muller, Bridgewater, NJ
ASSIGNEE: Celgene Corporation, Warren, NJ (U.S. corp.)

08/703,708 APPL-NO: DATE FILED: Aug. 27, 1996 ART-UNIT: 123

PRIM-EXMR:

Alan L. Rotman ASST-EXMR: D. Margaret M. Mach

LEGAL-REP: Mathews, Collins, Shepherd & Gould, P.A.

US PAT NO: 5,698,579 [IMAGE AVAILABLE] US-CL-CURRENT: **514/416**; 548/512

L15: 47 of 72

SUMMARY:

BSUM(12)

The **nuclear** **factor** .**kappa**.**B** (NF.**kappa**.B) is a pleiotropic transcriptional activator (Lenardo, et al. Cell 1989, 58, 227-29). NF.kappa.B has been implicated as a transcriptional activator.

US PAT NO: 5,691,338 [IMAGE AVAILABLE] DATE ISSUED: Nov. 25, 1997

L15: 48 of 72

1,2-dithiole-3 thiones for the treatment of reverse

transcriptase-dependent viral infections R: Hans J. Prochaska, New York, NY

INVENTOR:

Bruce Polsky, New York, NY

ASSIGNEE: Sloan-Kettering Institute for Cancer Research, New York,

NY (U.S. corp.) 08/485,658

APPL-NO: DATE FILED: Jun. 7, 1995

ART-UNIT: 124

PRIM-EXMR: Samuel Barts

John P. White LEGAL-REP:

US PAT NO: 5,691,338 [IMAGE AVAILABLE]
US-CL-CURRENT: **514/252**, **262**, **274**, **441* 1.15: 48 of 72

SUMMARY:

BSUM(5)

This . . . by which these thiols inhibit HIV-1 replication is believed to be due to their ability to inhibit the activation of
"Nuclear*" **Factor*" . **kappa** **B** under conditions of oxidative stress (7). There has been enthusiasm for testing compounds such as N-acetylcysteine or the ester of. . .

DETDESC:

7. Staal F. J., M. Roederer, and L. A. Herzenberg. Intracellular thiols regulate activation of **nuclear** **factor** **kappa** **B** and transcription of human immunodeficiency virus. Proc. Natl. Acad. Sci. 87:9943-9947 (1990).

US PAT NO: 5,686,436 [IMAGE AVAILABLE]

L15: 49 of 72

DATE ISSUED: Nov. 11, 1997

Multi-faceted method to repress reproduction of latent viruses in humans and animals TITLE:

INVENTOR: Knox Van Dyke, Morgantown, WV

ASSIGNEE: HIV Diagnostics, Inc., Lexington, KY (U.S. corp.) 08/317,730 APPL-NO:

DATE FILED: Oct. 4, 1994

ART-UNIT: 152

PRIM-EXMR:

Gollamudi S. Kishore

LEGAL-REP: Price, Heneveld, Cooper, DeWitt & Litton

US PAT NO: 5,686,436 [IMAGE AVAILABLE] L15: 49 of 72 US-CL-CURRENT: **514/171**, **198**, **369**, **374**, **378**, **561**, **563**

such as HIV, in animals by the generally concurrent Disclosed . . administration of (1) antioxidants including a glutathione agent; and (2) an **NFKB** induction inhibitor. Also disclosed are pharmaceutical compositions and kits for use in repressing reproduction of latent viruses such as HIV.

SUMMARY:

BSUM(8)

Schreck . . . The iKB factor is removed from the protein triad and the remaining p50, p65 complex becomes known as NF-kappa B (**NFKB**).

SUMMARY:

BSUM(9)

Schreck et al. have recognized that **NFKB** is a gene transcription factor that migrates into the nucleus of the HIV infected cell and switches on the production. . . expression of HIV-1 in a human T cell line. They further report that the expression of HIV is mediated by **NFKB** transcription factor which is potently and rapidly activated by a hydrogen peroxide treatment of cells from its inactive cytoplasmic form. They additionally report that N-acetyl cysteine and other thiol compounds block the activation of "NFKB". They concluded that these diverse agents thought to activate "NFKB" by distinct intracellular pathways might act through a common mechanism involving the synthesis of reactive oxygen intermediates. They did not. . .

SUMMARY:

BSUM(10)

Sherman et al., Biochem. Biophys. Res. Comm., 191 (3):1301-1308, 1993, report that pyrrolidine dithiocarbamate (PDTC) is an inhibitor of **NFKB** activation. They further report that this compound is an inhibitor of nitric oxide synthase (NO synthase). They further report that. . . that PDTC may act as a scavenger of reactive oxygen species which prevents them from participation in the activation of **NFKB**.

SUMMARY:

BSUM(15)

The . . . the generally concurrent administration of 1) a glutathione agent; 2) at least one additional antioxidant; and 3) at least one **NFKB** induction inhibitor. Further aspects and advantages of the invention will be apparent to those skilled in the art upon review. . .

DETDESC:

DETD(3)

There . . . glutathione precursor, a glutathione production enhancer, or glutathione, (2) high doses of additional fat- and water-soluble antioxidants, and (3) an **NFKB** induction inhibitor, to an animal infected with a latent virus. The fat- and water-soluble antioxidants are administered to an animal. . .

DETDESC:

DETD(5)

The Role of **NFKB** and Peroxynitrite in the Activation of a Cell to Reproduce HIV

DETDESC:

DETD(6)

NFKB is a gene transcription factor that switches on the production of the HIV virus of a vitally infected cell. **NFKB** is known to activate a variety of genes, including the transcription of a variety of cytokines, viruses and NO Synthase.. . .

DETDESC:

DETD(10)

Peroxynitrite is significant in that it activates **NFKB** **NFKB** is inactivated by I Kappa B (IKB) which acts on **NFKB** via the P65 subunit. As shown in FIG. 1, peroxynitrite cleaves IKB, thereby releasing the active **NFKB**

DETDESC:

DETD(28)

NFKB Induction Inhibitors

DETDESC:

DETD(29)

NFKB induction inhibitors are agents that inhibit **NFKB** transcription factor from binding to DNA. This blocks the induction of HIV or other viral reproduction by directly suppressing the vital reproduction activating mechanism. **NFKB** inhibitors (item 7, FIG. 2) also suppress peroxynitrite synthesis, by preventing **NFKB** from activating cell genes to produce NO synthase.

DETDESC:

DETD(32)

The preferred type of **NFKB** induction inhibitor is an anti-inflammatory steroid. Examples of suitable anti-inflammatory steroids suitable as **NFKB** induction inhibitors include but are not limited to predonsone, prednisolone, methyl prednisolone, dexamethasone, beta metasone dehydroepiandrosterone, 9a-fluorocortisol, prednisone, aetiocholanolone, 2-methyltoortisol, pregnanediol, deoxycorticosterone, cortisone, hydrocortisone (cortisol), 6a-methylprednisolone, triamcinolone, estrogen or derivatives thereof. Generally, any steroid with antitinflammatory action toward **NFKB** may be used. In addition, one or more suitable nongluocoorticoid lazaroids may be utilized as **NFKB** induction inhibitors. Preferred lazaroids include, but are not limited to, U-74006F, which is 21-[4-(2,6-di-1-pyrrolidinyl-4-pyrimidinyl)-1-piperazinyl]-16-methyl-(16.alpha.)-pregna-1,4,9(11)-triene-3,20-dione monomethanesulfonate or TIRLLAZAD mesylate or.

DETDESC:

DETD(36)

In . . . Other antiinflammatory steroids can be substituted at appropriate doses, as set forth in the Physicians' Desk Reference. Administration of an **NFKB** induction inhibitor such as an anti-inflammatory steroid, is one of the most important steps in the treatment of HIV. AIDS.

DETDESC:

DETD(39)

In addition, to the previously noted anti-inflammatory steroids and lazaroids, a variety of other compounds may be utilized as ""NFKB" induction inhibitors such as pyrrolidine dithiocarbamate and other dithiocarbamates, and glycyrrhizic acid (from licorice root). A preferred dosage level when. . . is about 100 mg/day per person for each day of therapy. In addition, other compounds are suitable for use as ""NFKB" induction inhibitors. These inhibitors include, but are not limited to, immunosuppressants such as cyclosporin A, rapamycin, interleukin 10, and FK. . . Clearly, a wide array of plant steroids, male steroids, female steroids, glucocorticoids, lazaroids, and 21-aminosteroids are eligible for use as ""NFKB" induction inhibitors.

DETDESC:

DETD(40)

An inhibitor known to be effective against **NFKB** binding or expressing is mevinolin, a drug which prevents isoprenylation and methylthioadenosine (MTA) and inhibitor of several S adenosylmethionine dependent.

DETDESC:

DETD(47)

Although . . . antioxidants, glutathione agents, and steroids with regard to HIV production. HIV replication is blocked by a combination of antioxidants and "NFKB" induction inhibitor. About 70% of the blocking action of HIV replication is believed to stem from the "NFKB" induction inhibitor, which preferably is one or more anti-inflammatory steroids. Although such steroids do not have direct inhibitory activity, they control viral synthesis by blocking "NFKB" induction. As will be recalled, "NFKB" is a DNA transcription factor made of protein. "NFKB" controls a whole series of inflammatory cytokines and NO synthase as well as HIV and FIV replication. Upon introduction of. . .

DETDESC:

DETD(48)

However, for **NFKB** to be active it must shed its inhibitory factor I kappa B. Such shedding requires oxidation because the bonds holding. . to proteins F90 and P65 are sensitive to oxidation. Thus, antioxidants keep the I kappa B inhibitory factor bound to **NFKB** and therefore inactive.. The role of antioxidants in the mechanism depicted in FIG. 3 is believed to be responsible for about 30% of the activity of producing **NFKB** and preventing HIV replication.

DETDESC:

DETD(49)

All... known to those skilled in the art. Although it is most preferred to administer the anitoxidants including glutathione agent and "NFKB" induction inhibitor concurrently, or simultaneously, it is not a requirement. Thus, the preferred embodiments of the present invention also encompass.

DETDESC:

DETD(51)

The . . . a glutathione agent; (2) an effective amount of one or more additional antioxidants; and (3) an effective amount of an **NFKB**

induction inhibitor. In a most preferred embodiment, the pharmaceutical compositions comprise: (1) an effective amount of a glutathione agent, e.g.. . antioxidant, (2b) an effective amount of a fat-soluble antioxidant, and (3) an effective amount of an anti-inflammatory steroid as the "NFKB" induction inhibitor. The other ingredients described above may also be included.

DETDESC:

DETD(58)

In . . . C, A and E; an effective amount of at least one glutathione precursor such as N-acetyl cysteine; followed by an **NFKB** induction inhibitor such as one or more anti-inflammatory steroids or lazaroids. As summarized in Table 4 below, seven cats heavily. . . 10 to about 18 pounds. The cats were initially treated with a single dosage of an effective amount of an **NFKB** induction inhibitor, that is an antiinflammatory steroid dose of DEPO-MEDROL (20-25 mg) and a series of oral dosages of a. . .

DETDESC:

DETD(62)

In . . . fat-soluble antioxidants and an effective amount of at least one glutathione precursor such as N-acetyl cysteine are administered, before an **NFKB** induction inhibitor is administered, the CD.sub.4 (T-lymphocyte) count is increased to about 100 cells/mm.sup.3 or more. The CD.sub.4 count may. . . concentrates containing monocytes may be given, such as via transfusions. Once CD.sub.4 counts are about 100 cells/mm.sup.3 or more, an **NFKB** induction inhibitor is administered.

DETDESC:

DETD(63)

In both the preferred and optional treatment regimens, the **NFKB** induction inhibitor is administered until AIDS(-) is indicated from AIDS(+) blood assay, via ELISA Western blot, and PCR (polymerase chain.

DETDESC:

DETD(74)

Preferably, . . . suitable glutathione precursors could be utilized in place of, or instead of the N-acetyl cysteine. Similarly, one or more other "*NFKB*" induction inhibitors could be utilized in place of or instead of the methyl prednisolone.

DETDESC:

DETD(77)

The . . . one fat soluble antioxidant at doses higher than the recommended daily minimum requirements, and preferably, only slight amounts or no **NFKB** induction inhibitor. In a most preferred treatment regimen, the subject suffering from symptoms of the Herpes virus is administered generally. . .

CLAIMS:

CLMS(1)

The . . .

for suppressing the reproduction of human immunodeficiency virus in a human, comprising administering to such human:
(i) at least one **NFKB** induction inhibitor in an amount effective to inhibit **nuclear** **factor** **kappa** **B**; said at least one **NFKB** induction inhibitor being selected from the group consisting of anti-inflammatory steroids and nonglucocorticoid lazaroids;
(ii) at least one fat-soluble antioxidant at. . .

CLAIMS:

CLMS(13)

 The method of claim 1 wherein said **NFKB** induction inhibitor comprises an anti-inflammatory steroid.

CLAIMS:

CLMS(14)

14.... 1 further comprising administering to such human:
 (v) an effective amount of a peroxynitrite production suppressor in addition to said **NFKB** induction inhibitor.

CLAIMS

CLMS(15)

15. . . . suppressing the reproduction of feline immunodeficiency virus and/or feline leukemia virus, comprising administering to a cat:

(i) at least one **NFKB** induction inhibitor in art amount effective to inhibit **nuclear** **factor** **kappa** **B**; said at least one **NFKB** induction inhibitor being selected from the group consisting of anti-inflammatory steroids and nonglucocorticoid lazaroids;
(ii) at least one fat-soluble antioxidant at. . .

CLAIMS:

CLMS(20)

20. The method of claim 15 wherein said **NFKB** induction inhibitor comprises an anti-inflammatory steroid.

CLAIMS:

CLMS(21)

21. . . . 15 further comprising administering to such cat: (v) an effective amount of a peroxynitrite production suppressor in addition to said **NFKB** induction inhibitor.

CLAIMS:

CLMS(26)

26. . . . 24 further comprising administering to such animal: (v) an effective mount of a peroxynitrite production suppressor in addition to said **NFKB** induction inhibitor.

CLAIMS:

CLMS(30)

30. . . . 28 further comprising administering to such animal; (v) an effective mount of a peroxynitrite production suppressor in addition to said **NFKB** induction inhibitor.

US PAT NO: 5,683,987 [IMAGE AVAILABLE]

L15: 50 of 72

L15: 50 of 72

DATE ISSUED: Nov. 4, 1997 TITLE:

Therapeutic oligonucleotides targeting the human MDR1 and MRP genes

INVENTOR: Larry J. Smith, Omaha, NE

ASSIGNEE: The Board of Regents of the University of Nebraska,

Lincoln, NE (U.S. corp.) 08/487,141

APPL-NO:

DATE FILED: Jun. 7, 1995 189

ART-UNIT:

George G. Elliott PRIM-FYMR-

ASST-EXMR: Andrew Wang
LEGAL-REP: Dann, Dorfman, Herrell and Skillman

US PAT NO: 5,683,987 [IMAGE AVAILABLE]
US-CL-CURRENT: **514/44**; 536/23.1, 24.31, 24.5

SUMMARY:

BSUM(10)

In . . . transplant models (Kitajima et al., J. Biol. Chem. 267:25881-25888, 1992). Others have targeted genes in cancer cells, including c-myc, c-Ha-ras, **NF**-**kB**, c-myb, c-kit and bcr-abl. In each of these instances involving the administration of ODNs to treat animals with xenogeneic human. . .

DETDESC:

DETD(62)

556 22 cap site

LOW(3)mdr 20 low Tm +

Cohen(1)mdr

1130 15 published

NF-**kB**(1)mdr 296 22 3; TR --

binding

CAT(L)mdr 432 20 TR binding

Y-box-mdr 464 22 TR binding

US PAT NO: 5,679,684 [IMAGE AVAILABLE]
DATE ISSUED: Oct. 21, 1997

1.15: 51 of 72

Xiannong Chen, Athens, GA George J. Cianciolo, Chapel Hill, NC Jose-Luis Diaz, Durham, NC

Bradley J. Benson, Chapel Hill, NC

Hydroxyalkylammonium-pyrimidines and nucleoside derivatives, useful as inhibitors of inflammatory

Khalid S. Ishaq, Chapel Hill, NC Susan L. Morris-Natschke, Apex, NC Ronald J. Uhing, Durham, NC

Henry Wong, Morrisville, NC

ASSIGNEE: Macronex, Inc., Morrisville, NC (U.S. corp.)

The University of N.C. at Chapel Hill, Morrisville, NC

(U.S. corp.) APPL-NO: 08/476,704 DATE FILED: Jun. 7, 1995 ART-UNIT: 122

cytokines

PRIM-EXMR: Yogendra N. Gupta Klauber & Jackson LEGAL-REP:

US PAT NO: 5,679,684 [IMAGE AVAILABLE] US-CL-CURRENT: **514/269**, **50**, **274** L15: 51 of 72

SUMMARY:

BSHM(3)

TITLE:

INVENTOR:

Two . . . that they employ different signal transduction pathways. While both receptors are capable of binding TNF and activating the transcription factor **NFkB**, it appears that the expression of each receptor is independently and differentially regulated. Human TNF-.alpha. will bind to both types. . .

US PAT NO: 5,663,153 [IMAGE AVAILABLE] DATE ISSUED: Sep. 2, 1997

L15: 52 of 72

TITLE: Immune stimulation by phosphorothioate oligonucleotide analogs

INVENTOR: Stephen L. Hutcherson, Richmond, VA Josephine M. Glover, Woking, United Kingdom

ASSIGNEE: Isis Pharmaceuticals, Inc., Carlsbad, CA (U.S. corp.)

APPL-NO: 08/467,930 DATE FILED: Jun. 6, 1995

ART-UNIT: 189

PRIM-EXMR: Charles C.P. Rories

LEGAL-REP: Law Offices of Jane Massey Licata

US PAT NO: 5,663,153 [IMAGE AVAILABLE] L15: 52 of 72 US-CL-CURRENT: **514/44**; 424/1.11, 1.73, 1.77, 278.1, 280.1; 536/23.1, 24.5

SUMMARY:

BSUM(18)

Oligonucleotides having a sequence identical to a portion of the sense strand of the mRNA encoding the p65 subunit of **NF**-**kB**, a DNA binding protein, were found to stimulate splenic cell proliferation both in vitro and in vivo. The proliferating spleen cells were shown to be B cells. Immunoglobulin secretion and **NF**.**kB** activity in these cell lines was also increased by the sense oligonucleotide. Both phosphodiester and phosphorothioate sense oligonucleotides stimulated

US PAT NO: 5,658,940 [IMAGE AVAILABLE] DATE ISSUED: Aug. 19, 1997

TITLE: Succinimide and maleimide cytokine inhibitors
INVENTOR: George W. Muller, Bridgewater, NJ
Mary Shire, North Plainfield, NJ

ASSIGNEE: Celgene Corporation, Warren, NJ (U.S. corp.)

APPL-NO: 08/539.879 DATE FILED: Oct. 6, 1995

ART-UNIT: 121

PRIM-EXMR: Jacqueline Haley

LEGAL-REP: Mathews, Collins, Shepherd & Gould, P.A.

US PAT NO: 5,658,940 [IMAGE AVAILABLE] L15: 53 of 72 US-CL-CURRENT: **514/417**, **309**, **339**, **340**, **421**, **425**; 546/142, 277.1, 278.4, 278.7; 548/465, 479, 513, 547

SUMMARY:

BSUM(14)

The **nuclear** **factor** .**kappa**.**B** (NF.**kappa**.B) is a pleiotropic transcriptional activator (Lenardo, et al. Cell 1989, 58, 227-29), NF.kappa.B has been implicated as a transcriptional activator.

US PAT NO: 5,650,316 [IMAGE AVAILABLE] DATE ISSUED: Jul. 22, 1997 L15: 54 of 72 TITLE: Uses of triplex forming oligonucleotides for the treatment of human diseases INVENTOR: Bharat B. Aggarwal, Houston, TX Robert F, Rando, The Woodlands, TX
Michael E, Hogan, The Woodlands, TX
E: Research Development Foundation, Carson City, NV (U.S. ASSIGNEE: corp.) APPL-NO: 08/254,114 DATE FILED: Jun. 6, 1994 ART-UNIT: 189 PRIM-EXMR: Charles C. P. Rories LEGAL-REP: Benjamin Aaron Adler US PAT NO: 5,650,316 [IMAGE AVAILABLE] L15: 54 of 72 US-CL-CURRENT: 435/375, 6, 7.23; **514/44**; 536/24.31, 24.32, 24.33, 24.5 DRAWING DESC: DRWD(11) FIG. 9 shows the antiproliferative effects of the TFOs J111-50 (Intron 3) and J109-50 (**NF**-**kB**) on a human glioblastoma (U-251) cell line. DETDESC DETD(9) 990 cholesterol scramble sequence of J111-51 B106-96 phosphorothioate G-rich TFO 1208 phosphorothioate random cholesterol TNF (**NF**-**kB**); -237 to -238 1109-51 DETDESC: DETD(53) FIGS. 9, top and bottom the antiproliferative effects of the TFOs J111-50 (Intron 3) and J109-50 (**NF**.**kB**) on a human glioblastoma (1.251) cell line, respectively. Both J111-50 (directed to Intron 3) and J109-50 (directed to Intron 3) and J109-50 (directed to Intro 3) and I109-50 (directed to Intro 3) and I11-50 (directed t DETDESC: DETD(63) Table . . . untreated cells was expressed as 100%. All determinations were made in triplicate. TFO 109-50, 111-51, 108-56 and 108-57 are from **NF**-**KB** (-237 to -208); Intron 3 (+1429 to +1456); and SP-1 (-58 to -33) sites respectively. J109-50 had amino group whereas. . . US PAT NO: 5,646,185 [IMAGE AVAILABLE] DATE ISSUED: Jul. 8, 1997 L15: 55 of 72 INVENTOR: Tumor treatment method
INVENTOR: Amato J. Giaccia, Stanford, CA
Albert C. Koong, Palo Alto, CA
ASSIGNEE: The Board of Trustees of the Leland Stanford Junior University, Stanford, CA (U.S. corp.) APPL-NO: 08/137,238 DATE FILED: Oct. 14, 1993 ART-UNIT: 124 Paul J. Killos PRIM-EXMR: Susan T. Evans, Carol A. Stratford, Peter J. Dehlinger LEGAL-REP: US PAT NO: 5,646,185 [IMAGE AVAILABLE] L15: 55 of 72 US-CL-CURRENT: **514/548** DRAWING DESC: FIG. 6B shows sequences of PKC responsive elements **NFKB** (SEQ ID NO: 2), HSE (SEQ ID NO: 3), GRE SEQ ID NO: 4 and API-1 (SEQ ID NO: 5). . . DETDESC:

DETD(61)

As . . . upstream of the TNF gene. Exemplary PKC Responsive elements that may be included in the vector include Glucose-related Core element, **Nuclear** **Factor**-**kappa** **B** (**NFKB**), Heat shock

transcription factor (HSE), GRE and AP-1. Sequences for these exemplary elements are shown in FIG. 6B. Inclusion of. . .

US PAT NO: 5,643,893 [IMAGE AVAILABLE] DATE ISSUED: Jul. 1, 1997
TITLE: N-substituted-(Dihydroxyboryl)alkyl purine, indole and pyrimidine derivatives, useful as inhibitors of inflammatory cytokines
Bradley J. Benson, Chapel Hill, NC INVENTOR:

Xiannong Chen, Athens, GA George J. Cianciolo, Chapel Hill, NC Jose-Luis Diaz, Durham, NC Khalid S. Ishaq, Chapel Hill, NC Susan L. Morris-Natschke, Apex, NC Ronald J. Uhing, Durham, NC Henry Wong, Morrisville, NC

E: Macronex, Inc., Wayne, PA (U.S. corp.)
University of North Carolina, Chapel Hill, NC (U.S. corp.) ASSIGNEE:

APPL-NO: 08/264,039 DATE FILED: Jun. 22, 1994 ART-UNIT: 122
PRIM-EXMR: Emily Bernhardt LEGAL-REP: Klauber & Jackson

US PAT NO: 5,643,893 [IMAGE AVAILABLE] US-CL-CURRENT: **514/64**; 544/229; 548/405; 562/7 L15: 56 of 72

SHMMARY.

BSUM(3)

. that they employ different signal transduction pathways. While both receptors are capable of binding TNF and activating the transcription factor **NFkB**, it appears that the expression of each receptor is independently and differentially regulated. Human TNF-.alpha. will bind to both types.

US PAT NO: 5,641,773 [IMAGE AVAILABLE] DATE ISSUED: Jun. 24, 1997 L15: 57 of 72 Methods for treating viral infections

INVENTOR: DR: Arthur P. Pardee, Brookline, MA Debajit K. Biswas, Newton, MA Bruce J. Dezube, Newton Centre, MA

ASSIGNEE: Dana-Farber Cancer Institute, Boston, MA (U.S. corp.) 08/159,509

APPL-NO: DATE FILED: Nov. 30, 1993 ART-UNIT: 129 PRIM-EXMR: Bri Brian M. Burn

LEGAL-REP: David G. Conlin, Ronald I. Eisenstein

US PAT NO: 5,641,773 [IMAGE AVAILABLE] L: US-CL-CURRENT: **514/221**, **258**, **262**, **264** L15: 57 of 72

DETDESC:

DETD(4)

Mapping . . . in trans-activating (stimulating) viral gene expression by interaction with the tar element which is present in this region.

Similarly, the ""nuclear" "factor" NF- "kappa". "B" can also stimulate viral gene expression through its interaction with sequences present in the LTR [Sen, R., et al, Cell. . .

US PAT NO: 5,635,517 [IMAGE AVAILABLE] DATE ISSUED: Jun. 3, 1997

Method of reducing TNF alpha. levels with amino substituted 2-(2,6-dioxopiperidin-3-yl)-1-oxo-and 1,3-dioxoisoindolines TITLE:

INVENTOR: George W. Muller, Bridgewater, NJ

David I. Stirling, Branchburg, NJ

Roger S. -C. Chen, Edison, NJ

E: Celgene Corporation, Warren, NJ (U.S. corp.) ASSIGNEE:

08/690,258 APPL-NO: DATE FILED: Jul. 24, 1996 ART-UNIT: PRIM-EXMR: 123 C. Warren Ivy C. S. Aulakh

ASST-EXMR:

LEGAL-REP: Mathews, Collins, Shepherd & Gould, P.A

US PAT NO: 5,635,517 [IMAGE AVAILABLE] L15: 58 of 72 US-CL-CURRENT: **514/323**; 546/201

SUMMARY:

BSUM(11)

The **nuclear** **factor** .**kappa**.**B** (NF.**kappa**.B) is a

pleiotropic transcriptional activator (Lenardo, et al. Cell 1989, 58, 227-29). NF kappa. B has been implicated as a transcriptional activator.

US PAT NO: 5,629,152 [IMAGE AVAILABLE] DATE ISSUED: May 13, 1997

L15: 59 of 72

Trisubstituted .beta.-lactams and oligo

.beta.-lactamamides

Vasulinga Ravikumar, Carlsbad, CA INVENTOR: Isis Pharmaceuticals, Inc., Carlsbad, CA (U.S. corp.)

ASSIGNEE: 08/283,591

DATE FILED: Aug. 1, 1994

ART-UNIT: 187

PRIM-EXMR: Stephanie W. Zitomer

ASST-EXMR: Dianne Rees

Woodcock Washburn Kurtz MacKiewicz & Norris LEGAL-REP:

US PAT NO: 5,629,152 [IMAGE AVAILABLE] US-CL-CURRENT: 435/6, 91.1; **514/44**; 536/24.3, 24.5

L15: 59 of 72

DETDESC:

DETD(125)

Phosphorothioate . . . (Waters, Division of Millipore Corp., Milford, Ma.) and ethanol precipitated. The phosphorothioate oligonucleotides are hybridized to create the double stranded **NF**-**kB** binding site.

DETDESC:

DETD(129)

C-rel has been shown to represent a constituent of the **NF**-**kB** site binding transcription factor, which plays a crucial role in the expression of a number of genes including the immunoglobulin.

DETDESC:

DETD(130)

Crude . . . of poly dl.dC as a nonspecific competitor at a concentration of 100 .mu.g/ml of extract. Nuclear extracts containing the biotinylated **NF**-**kB** binding site competitor are prepared as in Example 34, above.

DETDESC:

DETD(131)

A series of oligo .beta.-lactamamide duplexes is synthesized to correspond to various length fragments of the consensus binding sequence of c-rel. **NF**-**kB** binding site competitor is added to each duplex and the resulting samples are washed. Antibody directed to rel is added..

US PAT NO: 5,624,912 [IMAGE AVAILABLE]
DATE ISSUED: Apr. 29, 1997

L15: 60 of 72

Method of treating HIV infection and related secondary

infections with defibrotide

OR: Arsinur Burcoglu, 213 Sweetgum Rd., Pittsburg, PA 15238 Marc Wagner, 4201 Greensburg Pike, Pittsburg, PA 15221 INVENTOR:

APPL-NO: 08/185.416

DATE FILED: Jan. 24, 1994

ART-UNIT: 189

PRIM-EXMR:

Deborah Crouch Banner & Witcoff, Ltd LEGAL-REP:

L15: 60 of 72

US PAT NO: 5,624,912 [IMAGE AVAILABLE] US-CL-CURRENT: **514/44**, **924**, **934**

DETDESC:

DETD(118)

The transcription factor **NF**-**kB** binds to both the HIV-I enhancer, and the sILR2 gene. Protein kinase C phosphorylates its inhibitor IkB and releases active **NF**-**kB**. Increased cAMP levels by inhibiting directly the Ca.sup.2+ induced activation of protein kinase C would modulate this phosphorylation event, and downregulate the transcriptional activities related to **NF**-**kB**. Since **NFkB** binds to both the HIV enhancer and IL2 receptor, increased cAMP levels will downregulate HIV-I replication.

DETDESC:

m) defibrotide sequence+LTR **NFkB** mutant (-104 to -80),

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US PAT NO: 5,612,330 [IMAGE AVAILABLE] DATE ISSUED: Mar. 18, 1997
```

OR: David T. Connor, Ann Arbor, MI Stephen J. Gracheck, Ann Arbor, MI

ASSIGNEE: Warner-Lambert Company, Morris Plains, NJ (U.S. corp.)

APPL-NO: 08/408,431

DATE FILED: Mar. 22, 1995

ART-UNIT: 122

PRIM-EXMR: Robert T. Bond

LEGAL-REP: Charles W. Ashbrook

US PAT NO: 5,612,330 [IMAGE AVAILABLE]

L15: 61 of 72

1.15: 61 of 72

US-CL-CURRENT: **514/211**, **220**; 540/495

SUMMARY:

BSUM(7)

An . . . further development is therapeutic targeting along cellular signaling pathways that result in HIV-1 transcriptional activation. Among the potential targets is **nuclear** **factor**. **kappa**. **B** (NF-, **kappa**.B), a transcriptional enhancer important for HIV-1 activation. In resting cells, preformed NF-, kappa. B exists in the cytoplasm bound to its inhibitor. .

US PAT NO: 5,605,914 [IMAGE AVAILABLE] DATE ISSUED: Feb. 25, 1997 TITLE: Imides

L15: 62 of 72

INVENTOR: George W. Muller, Bridgewater, NJ

ASSIGNEE: Celgene Corporation, Warren, NJ (U.S. corp.) 08/258.587

APPL-NO: DATE FILED: Jun. 10, 1994

ART-UNIT:

PRIM-EXMR: C. Warren Ivy

ASST-EXMR-

D. Margaret M. Mach Mathews, Woodbridge & Collins LEGAL-REP:

US PAT NO: 5,605,914 [IMAGE AVAILABLE] L15: 62 (US-CL-CURRENT: **514/339**, **417**; 546/277.1; 548/465, 479 1.15: 62 of 72

SUMMARY:

BSUM(14)

The **nuclear** **factor** kB (NF. **kappa**. **B**) is a pleiotropic transcriptional activator (Lenardo, et al. Cell 1989, 58, 227-29). NF, kappa, B has been implicated as a transcriptional activator. .

US PAT NO: 5,596,011 [IMAGE AVAILABLE] DATE ISSUED: Jan. 21, 1997

L15: 63 of 72

1.15: 64 of 72

Method for the treatment of macular degeneration INVENTOR: Karen M. Repine, 2275 Cherry Hills Farm Dr., Englewood, CO

80110

John E. Repine, 2275 Cherry Hills Farm Dr., Englewood, CO

80110

APPL-NO: 08/418,645

DATE FILED: Apr. 6, 1995 ART-UNIT: 125

PRIM-EXMR: Zohreh Fay

US PAT NO: 5.596.011 [IMAGE AVAILABLE] L15: 63 of 72 US-CL-CURRENT: **514/369**, **562**, **665**, **912**

DETDESC:

DETD(4)

. . protective mechanisms offered by N-acetylcysteine (NAC) include: scavenging (inactivation of) oxygen radicals (e.g. H.sub.2 O.sub.2, HOCl or .circle-solid.OH) directly; inhibiting **NFkB** nuclear factor activation, which is a known link between viral infection and activation of oxidative processes; decreasing oxidant-induced lipid peroxidation. .

US PAT NO: 5,583,155 [IMAGE AVAILABLE] DATE ISSUED: Dec. 10, 1996

6-amino-1,2-benzopyrones useful for treatment of viral TITLE:

diseases

INVENTOR: Ernest Kun, Mill Valley, CA Laure Aurelian, Baltimore, MD

Octamer, Inc., Mill Valley, CA (U.S. corp.)

ASSIGNEE: APPL-NO: 08/237,969

DATE FILED: May 3, 1994

125 ART-UNIT:

PRIM-EXMR: T. J. Criares

LEGAL-REP: Albert P. Halluin, Scott R.Pennie & Edmonds Bortner

US PAT NO: 5,583,155 [IMAGE AVAILABLE] US-CL-CURRENT: **514/457**, **456**

L15: 64 of 72

DETDESC:

DETD(42)

Expression . . . by phorbol esters and lectins as described in Science, 108:117 (1948); and Ibid, 230:850 (1986). This stimulation is mediated by **NF**-**kB**, a factor that regulates transcription and Expression . binds to the twice-repeated 11-bp kB motif in the HIV enhancer Nature, 326, 711 (1987). Mutations within this site that eliminate the binding of **NF**- **kB** also abolish the increase in HIV gene expression in activated T cells. However, HIV persists in macrophages in which it.

DETDESC:

DETD(43)

AIDS . . . synthesized immediately after infection resulting in immediate-early gene products. These include the trans-activating genes that activate HIV-LTRcat involving induction of **NF**-**kB** activity. CMV also activates HIV expression. This is mediated by the CMV IE gene. However, it does not appear to require **NF**-**kB** activity.

US PAT NO: 5,571,797 [IMAGE AVAILABLE]

DATE ISSUED: Nov. 5, 1996

TITLE: Method of inducing gene expression by ionizing radiation INVENTOR: Tsuneya Ohno, Boston, MA

Ralph R. Weichselbaum, Chicago, IL

Donald W. Kufe, Wellesley, MA

Arch Development Corporation, Chicago, IL (U.S. corp.) ASSIGNEE:

08/241,863 APPL-NO: DATE FILED: May 11, 1994

PRIM-EXMR: Br PRIM-EXMR: Bruce R. Campell LEGAL-REP: Arnold White & Durkee

US PAT NO: 5,571,797 [IMAGE AVAILABLE] L15: 65 of 72 US-CL-CURRENT: **514/44**; 424/1.11, 1.49, 1.61, 1.65, 1.69, 93.2, 93.21, 450; 435/69.1, 69.5, 320.1; 536/24.1

DETDESC:

DETD(79)

Transcription . . . domains are well known in the art. Exemplary transcription factors having activation domains are GALA, c-Jun, viral protein VP-16, and **nuclear** **factor** NF-.**kappa**.**B**.

DETDESC:

DETD(94)

Nuclear **factor** NF-.**kappa**.**B** is a transcription factor. The activation domain of NF-.kappa.B comprises amino acid residue sequences from about residue position 414 to. . .

DETDESC:

DETD(270)

The . . . al., 1990; Hallahan, et al, 1991). Other studies have demonstrated that x-rays induce expression and DNA binding activity of the **nuclear** **factor** .**kappa**. **B** (NF-. **kappa**. B; Brach, et al., 1991).

DETDESC:

DETD(313)

Ionizing . . . which code for transcription factors. Other studies have demonstrated that ionizing radiation induces expression and DNA binding activity of the ""nuclear" "factor" .**kappa" "B" (NF. .**kappa".B). The activation of transcription factors likely represents a critical control point in transducing early nuclear signals to longer term changes. .

DETDESC:

DETD(328)

. phorbol ester-induced activation of the HIV-1 long terminal repeat. This antioxidant has also been found to inhibit activation of the ""nuclear" ""factor" ""kappa" ""B" (NF-. ""kappa" ". B) by phorbol esters and other agents such as H.sub.2 O.sub.2. The available findings

suggest that ROIs activate NF-, kappa. B by induced. . .

US PAT NO: 5,550,132 [IMAGE AVAILABLE]

L15: 66 of 72

DATE ISSUED: Aug. 27, 1996

Hydroxyalkylammonium-pyrimidines or purines and nucleoside TITLE: derivatives, useful as inhibitors of inflammatory

cytokines

Bradley J. Benson, Chapel Hill, NC INVENTOR:

Xiannong Chen, Athens, GA George J. Cianciolo, Chapel Hill, NC

Jose-Luis Diaz, Durham, NC

Khalid S. Ishaq, Chapel Hill, NC Susan L. Morris-Natschke, Apex, NC

Ronald J. Uhing, Durham, NC

Henry Wong, Morrisville, NC
E: University of North Carolina, Chapel Hill, NC (U.S. corp.)
Macronex, Inc., Morrisville, NC (U.S. corp.) ASSIGNEE:

APPL-NO:

08/264,026 DATE FILED: Jun. 22, 1994

ART-UNIT: 122

PRIM-EXMR: Yogendra N. Gupta

LEGAL-REP: Klauber & Jackson

5,550,132 [IMAGE AVAILABLE] L15: 66 of 72 US-CL-CURRENT: **514/269**, **274**; 544/311, 312, 313, 314

SUMMARY:

BSUM(3)

Two . . . that they employ different signal transduction pathways. While both receptors are capable of binding TNF and activating the transcription factor **NFkB**, it appears that the expression of each receptor is independently and differentially regulated. Human TNF-.alpha. will bind to both types. .

L15: 67 of 72

US PAT NO: 5,547,979 [IMAGE AVAILABLE] L1:
DATE ISSUED: Aug. 20, 1996
TITLE: TNF inhibition
INVENTOR: Siegfried B. Christensen, IV, Philadelphia, PA
Klaus M. Esser, Downingtown, PA

Philip L. Simon, Randolph, NJ

ASSIGNEE: SmithKline Beecham, Philadelphia, PA (U.S. corp.)

APPL-NO: 08/424,944 DATE FILED: Apr. 19, 1995

ART-UNIT: 121

PRIM-EXMR: David B. Springer

LEGAL-REP: Dara L. Dinner, Stephen Venetianer, Edward T. Lentz

US PAT NO: 5,547,979 [IMAGE AVAILABLE] US-CL-CURRENT: **514/424**; 548/550, 551

DETDESC:

DETD(29)

There . . . (1989)]. A molecular mechanism for the virus inducing activity is suggested by TNFs ability to activate a gene regulatory roteins (**NF**-**kB**) found in the cytoplasm of cells, which promotes HIV replication through binding to a viral regulatory gene sequence (LTR) (See., .

US PAT NO: 5,420,154 [IMAGE AVAILABLE]
DATE ISSUED: May 30, 1995
TITLE: TNF inhibitors

INVENTOR: Siegfried B. Christensen, IV, Philadelphia, PA

Klaus M. Esser, Downingtown, PA Philip L. Simon, Randolph, NJ

ASSIGNEE: SmithKline Beecham Corp., Philadelphia, PA (U.S. corp.) 07/852.180

APPL-NO: DATE FILED: Mar. 30, 1992

121 ART-UNIT:

PRIM-EXMR: David B. Springer

Dara L. Dinner, Stephen Venetianer, Edward T. Lentz LEGAL-REP:

US PAT NO: 5,420,154 [IMAGE AVAILABLE]

US-CL-CURRENT: **514/424**; 548/551

L15: 68 of 72

L15: 68 of 72

DETDESC:

DETD(29)

There . . . (1989)]. A molecular mechanism for the virus inducing activity is suggested by TNFs ability to activate a gene regulatory protein (**NF**-**kB**) found in the cytoplasm of cells, which prom HIV replication through binding to a viral regulatory gene sequence (LTR) (Sec..

L15: 69 of 72

US PAT NO: 5,380,747 [IMAGE AVAILABLE] DATE ISSUED: Jan. 10, 1995 TITLE:

Treatment for atherosclerosis and other cardiovascular and

inflammatory diseases R: Russell M. Medford, Atlanta, GA INVENTOR: Margaret K. Offermann, Atlanta, GA R. Wayne Alexander, Atlanta, GA

Emory University, Atlanta, GA (U.S. corp.) 07/969,934 ASSIGNEE:

APPL-NO: DATE FILED: Oct. 30, 1992

PRIM-EXMR: Ma Marianne M. Cintins ASST-EXMR: William R. Jarvis LEGAL-REP: Kilpatrick & Cody

US PAT NO: 5,380,747 [IMAGE AVAILABLE] L15: 69 of 72 US-CL-CURRENT: **514/423**, **210**, **212**, **315**, **476**, **477**

DETDESC:

DETD(28)

At the molecular level, PDTC has been shown to inhibit the activation of the transcriptional regulatory factor **Nf**-**kB*** in response to certain cytokine and non-cytokine stimuli (Schreck, Rieber et al. 1991; Schreck, Meier et al. 1992). However, by. . . has been discovered that endothelial cells activate VCAM-1 gene expression through an apparently novel transcriptional regulatory factor that is not **Nf**-**kB**. This suggests that PDTC may regulate endothelial cell gene expression through its effect on a new transcriptional regulatory protein. It. . .

US PAT NO: 5,317,019 [IMAGE AVAILABLE]
DATE ISSUED: May 31, 1994

L15: 70 of 72

TITLE:

Inhibition of interleukin-1 and tumor necrosis factor

production by monocytes and/or macrophages

INVENTOR: Paul E. Bender, Cherry Hill, NJ Don E. Griswold, North Wales, PA

Nabil Hanna, Solana Beach, CA

John C. Lee, Radnor, PA

Bartholomew J. Votta, Pottstown, PA Philip L. Simon, Randolph, NJ

Alison M. Badger, Bryn Mawr, PA

Klaus M. Esser, Downingtown, PA

ASSIGNEE: SmithKline Beecham Corp., Phildelphia, PA (U.S. corp.) 07/809,484 APPL-NO:

DATE FILED: Dec. 12, 1991

ART-UNIT: 123

C. Warren Ivy PRIM-EXMR:

ASST-EXMR: Raymond Covington

LEGAL-REP: Dara L. Dinner, Stephen Venetianer, Edward T. Lentz

US PAT NO: 5,317,019 [IMAGE AVAILABLE] L15: 70 of 72 US-CL-CURRENT: **514/224.2**, **230.5**, **258**, **303**, **338**, **338**, **339**

SUMMARY:

BSUM(366)

There . . . (1989)]. A molecular mechanism for the virus inducing activity is suggested by TNFs ability to activate a gene regulatory protein (**NF**-**kB**) found in the cytoplasm of cells, which promotes HIV replication through binding to a viral regulatory gene sequence (LTR) [See,. .

US PAT NO: 5,306,724 [IMAGE AVAILABLE]
DATE ISSUED: Apr. 26, 1994

L15: 71 of 72

Method for preventing and treating atherosclerosis

TITLE: INVENTOR: Dennis I. Goldberg, Palatine, IL

Clintec Nutrition Company, Deerfield, IL (U.S. corp.) ASSIGNEE:

APPL-NO: 07/930.183

DATE FILED: Aug. 17, 1992 125

ART-UNIT:

PRIM-EXMR: Frederick E. Waddell ASST-EXMR: William R. A. Jarvis Hill, Steadman & Simpson LEGAL-REP:

L15: 71 of 72

US PAT NO: 5,306,724 [IMAGE AVAILABLE] US-CL-CURRENT: **514/369**, **824**

SUMMARY:

BSUM(30)

Adhesion . . . and growth factors which exacerbate the injury. The inflammatory process is promoted by cytokine activation of nuclear

transcription factor kB (**NF**-**kB**). This transcription factor is known to control the expression of a number of genes that code for cytokines and other. . .

SUMMARY:

BSUM(32)

Intracellular free radicals and hydrogen peroxides may serve as second messengers, transducing the cytokine signal to activate **NF**-**kB** The activation of **NF**.**RB** by a variety of pro-inflammatory cytokines, including interleukin-1. Lipopolysaccharide, lectin, TNF-alpha., phorbol ester and calcium ionophore, can be blocked by thiol containing compounds. Elevation of intracellular glutathione levels has been demonstrated to prevent the induction of HIV replication by been definitional to prevent the induction of 111 replication by "*NF".**kB**. The inventor believes that this data supports the hypothesis that maintaining intracellular glutathione levels may prevent the induction of proinflammatory. . .

US PAT NO: 5,294,630 [IMAGE AVAILABLE]

DATE ISSUED: Mar. 15, 1994
TITLE: Treatment of inflammatory bowel disease
INVENTOR: David Blake, Droitwich, United Kingo David Blake, Droitwich, United Kingdom

Peter P. K. Ho, Carmel, IN Jill A. Panetta, Zionsville, 1N

David Rampton, London, United Kingdom Nicola Simmonds, London, United Kingdom

ASSIGNEE: Eli Lilly and Company, Indianapolis, IN (U.S. corp.)

London Hospital Medical College, London, England (foreign

L15: 72 of 72

corp.) 07/909,852 APPL-NO:

DATE FILED: Jul. 7, 1992

ART-UNIT: 125

PRIM-EXMR: Leonard Schenkman

Joseph A. Jones, Leroy Whitaker LEGAL-REP:

L15: 72 of 72

US PAT NO: 5,294,630 [IMAGE AVAILABLE] L15: 72 of US-CL-CURRENT: **514/372**, **378**, **380**, **403**, **404**

SUMMARY:

BSUM(8)

. inflammatory cascade which leads to IBD. Schreck, Reactive Oxygen Intermediates, as Apparently Widely Used Messengers in the Activation of the **NF**-**KB** transcription factor and HIV-1, EMBO Journal 10, 2247-58 (1991).